APPENDIX A

The following as of code accomplish two tasks:

- I) Calculation of the topomeric conformation for a particular molecule, assuming that the molecule is referenced by a particular row of a Tripos Molecular Spreadsheet (MSS). With minor adaptations this code could be used in other molecular modeling environments, such as Cerius 2, Quanta, or Insight.
- II) Calculation of the line stope assuming that the biological data and one or more columns of property data are stored in a Tripos Molecular Spreadsheet (MSS). Almost any other software for manipulating data in a spreadsheet or other tabular representation could be adapted to perform similar calculations, assuming a Tanimoto function for expressing "distances" between bitsets of equal cardinality.

Both sections of code include procedures written in two languages. The first is C, familiar to all programmers, and includes both all specialized structure declarations and also brief explanations of all functions used. The second is SPL, an interpretative language available within the SYBYL molecular modeling program, whose syntax is similar to a Unix shell script. The SPL language is described fully in the volume entitled SPL Manual, found within the documentation set for SYBYL 6.2, release date July 1995. This volume includes descriptions of all "expression generators" (functions returning a value) and "macro commands" not specifically explained below.

I. Topomeric Field Code:

A. SPL macro CHOM!BUILD3D. To build topomerically aligned 3D models, the third argument must have the value ALIGN, and the global associative array element CHOM!Align[ALICYC] must have the value All_trans. Code to allow user adjustment of these and other 3D model-building parameters appearing in this code as other elements of CHOM!Align[] is not shown.

- B. Under these circumstances the following SPL macro CHOM!Alltrans sets all torsions provided to their topomeric values.
- C. To determine the atoms defining each torsion to be adjusted, CHOM!Alltrans invokes the expression generator %trans_path(), which executes the following C subroutine SYB_MGEN_CONN_BEST, with its associated subroutines syb_mgen_conn_att_atoms,

get_path_mw, get_path_xyz, and (if debugging) ashow. No user-adjustable values are used by this code. All non-obvious include files and a brief functional description of subroutines external to this code are provided in section III below.

D. The computation of rotatable-bond-attenuated steric (and/or electrostatic, hydrogen bonding) fields for the topomerically aligned conformation is carried out by the C subroutine QSAR_FIELD_EVAL_RB_ATTEN, which uses the accompanying subroutine QSAR_FIELD_RB_WTS to generate an attenuated weight for each atom's contribution to the field(s). (Pseudo code for the latter subroutine appears in its header comment.) The attenuation factor (recommended value of 0.85) is a user-adjustable or "tailorable" value, here shown as COMFA!AGGREG_SCALING. The user-adjustable HBOND_RAD_SCALING parameter affects the steric "radius" of a hydrogen-bonding hydrogen.

II. Patterson-Distribution Validation Code

A. The SPL expression generator *lrt_fast* returns the slope of the "best" <u>line</u> along with the count of data points and the fractional area, within a "virtual" or conceptual graph of absolute differences in biological activities vs absolute differences in the diversity measurement to be validated. The format of its output appears in the header comment.

B. The short SPL expression generator *dochi* shows the computation of the chi-squared statistic resulting from the output of the lrt_fast expression generator.

C. The C code functions QSHELL_HIER_LRT, QSHELL_HIER_DO_LRT, and fpt_heapsort generate the results produced by lrt_fast. These routines generate the biological differences themselves but rely on some external procedure, not shown, to generate the distances between the diversity measurements. (The reason is that the method of calculating differences depends on the diversity parameter(s). Typically a Euclidean distance is calculated for scalar properties, or a Tanimoto difference is calculated for bitsets, and if multiple parameters are combined to form the diversity measurement to be validated then the relative weighting must also be specified by the user.)

Section III. Supporting information for interpretation of the C code in Sections I and II.

- A. Declarations of complex and non-standard data structures referenced by the declarations within these C procedures, specifically for molecules, atoms, and the regions, fields, and other user input information that are part of a CoMFA field description.
- B. Functional descriptions of all external subroutines called by these C procedures, ordered alphabetically.

```
# SECTION I-A. Macro BUILD_3D for generating and storing topomeric alignments
@macro BUILD_3D CHOM
# builds 3D models,
    storage in a database or in a conformer column
    either not-aligned (just uses Concord or as-is if from Unity,
        or minimizes input structure)
    or aligned for CoMFA (requires core structure as alignment template)
       with optional fixup of side chains, charge calculation
   $1 is row ids in current MSS
   $2 is storage code (will retrieve structure from same place or somewhere)
   $3 is align (U or A)
   $4 is basic building technique
# other arguments, used only if ALIGN is true, are elements
        of the global associative array CHOM!ALIGN
# set up mol retrieval from MSS to be fast and clean
localvar AFFECT_SUBSET_save
localvar EXAMINE_TAILOR_MODE_save
localvar HIGHLIGHT MSS_save
localvar INFORM_save
localvar INPUT_MODE_save
localvar RELATE save
localvar SHOW_MOLECULE_save
localvar USER_FUNCTION_save natmcore heavy ys
localvar align ma rid cgq_save tailor_bumps_save newc \
        a b max_save usehs rat yrat nrat noth
                               $TAILOR!EXAMINE!AFFECT_SUBSET
setvar AFFECT SUBSET_save
setvar EXAMINE_TAILOR_MODE_save $TAILOR!EXAMINE!EXAMINE_TAILOR_MODE
setvar HIGHLIGHT_MSS_save $TAILOR!EXAMINE!HIGHLIGHT_MSS
                               STAILOR! EXAMINE! INFORM
setvar INFORM_save
setvar INPUT_MODE_save
                               $TAILOR!EXAMINE!INPUT_MODE
                               $TAILOR! EXAMINE! RELATE
setvar RELATE save
                               $TAILOR!EXAMINE!SHOW_MOLECULE
setvar SHOW_MOLECULE_save
                               $TAILOR!EXAMINE!USER_FUNCTION
setvar USER_FUNCTION_save
setvar cgq_save $CGQ_TIMEOUT
set CGQ_timeout 0
setvar TAILOR!EXAMINE!AFFECT_SUBSET
                                               NONE
setvar TAILOR!EXAMINE!EXAMINE_TAILOR_MODE
                                               SILENT
                                               NO
setvar TAILOR!EXAMINE!HIGHLIGHT_MSS
                                               NO
setvar TAILOR! EXAMINE! INFORM
setvar TAILOR!EXAMINE!INPUT_MODE
                                               ROW COLUMN EXPR
                                               NO
setvar TAILOR!EXAMINE!RELATE
                                               YES
setvar TAILOR!EXAMINE!SHOW_MOLECULE
setvar TAILOR!EXAMINE!USER_FUNCTION
                                               NONE
setvar max save $TAILOR!MAXIMIN2!LS_STEP_SIZE $TAILOR!MAXIMIN2!MAXIMUM_ITERATION
setvar ma %table_attribute( MOL_AREA )
# if needed make new place to put output
setvar newc
switch %substr( $2 1 3 )
case NEW)
  setvar newc %math( %table( * COL COUNT ) + 1 )
```

```
table column sln %cat( CONF $newc )
case SYB)
  database open %qspr_table_db( %table_default() ) update
  table ATTRIBUTE SET CONFORMER 0
case )
  setvar newc %substr( $2 1 %math( %pos( _ $2 ) - 1 )
  TABLE CONFORMER $newc
endswitch
if %streql( %substr( $3 1 1 ) "A" )
# are we bump checking ?
  if $CHOM!Align[BUMPS]
     setvar tailor bumps save $TAILOR!GENERAL!bumps contact distance
     tailor set general bumps contact distance %math( $CHOM!Align[BUMPS] - 1.0 )
  endif
##
# STEP 1: prepare template fragment
##
  setvar mcore $CHOM!Align[ MCORE ]
# save original template
  setvar mcsav %molempty()
  copy $mcore $mcsav
  default $mcore >$nulldev
  if $CHOM!Align[DEBUG]
     label id *
   endif
   setvar capsln %cat( %sln( $mcore ) )
   setvar natcore %mol_info( $mcore NATOMS )
# IF the alignment template has just one free valence,
# make geometrically acceptable template by adding heavy atoms, minimizing
# else use as is
   setvar heavy TRUE
   fillvalence *-H* Hal >$nulldev
   if %gt( %math( %mol_info( $mcore NATOMS ) - $natcore ) 1 )
        copy $mcsav $mcore
        setvar heavy
   endif
   if $heavy
    for a in %atoms(<H*>-<H>)
      modify atom type $a C.3 >$nulldev
      modify atom name $a X1 >$nulldev
    endfor
   endif
   TAILOR SET MAXIMIN2 LS_STEP_SIZE 0.0001 MAXIMUM ITERATIONS 1000 | |
  MAXIMIN $mcore DONE INTERACTIVE >$nulldev
if $heavy
   for a in %atoms(X1)
     modify atom type $a HEV
                                >$nulldev
# must rename it !!
     modify atom name $a X1 >$nulldev
   endfor
   setvar ys %set create( %atoms(X1) )
# orient template so that an R points in the positive X direction
```

```
setvar rat %arg( 1 %set_unpack( $ys ) )
   setvar nrat %arg( 1 %atom info( $rat NEIGHBORS ) )
   setvar yrat %arg( 1 %set_unpack( %set_diff( \
        ORIENT USER $nrat $rat $yrat >$nulldev
endif
# identify all the non-primary atoms for FIT, in/out of the search pattern
# and all the basic torsions (bonds to Ys) that potentially need setting
   setvar tpat %arg( 1 %search2d( %cat( %sln( $mcore ) ) $capsln NoDup 0 y ) )
   setvar hvinpat
   setvar patats
   setvar tors
   setvar usehs
   setvar sybhvats %set create(%atoms(*-<H>))
   if %lt( %set_size( $sybhvats ) 3 )
        setvar usehs TRUE
        setvar sybhvats %set create(%atoms(*))
   endif
   for a in %range(1 %sln_atom_count( $capsln ) )
      if %or( "$usehs" "%not( %set_and( %sln_atom_symbol( $capsln $a ) \
                H,F,Cl,Br,I ) ) " )
# for FIT, need to know the SYBYL IDs of the heavy atoms
        setvar hvinpat $hvinpat $a
        setvar patats[ $a ] %sln_rgroup_sybid( $mcore $tpat $a )
        setvar patats[ $a ][ YS ] %set_and( "$ys" "%set_create( \
                %atom_info( $patats[ $a ] NEIGHBORS ) ) " )
# for each torsion root, need to save the SLN ID of an arbitrary
                heavy atom torsional definer
        if $patats[$a][YS]
setvar tors[ $a ] %set_and( %set_diff( "%set_create( \ %atom_info( $patats[ $a ] NEIGHBORS ) )" $patats[ $a ][ YS ] ) $sybhvats )
# if there are several possibilities, prefer the lowest #'d carbon
                        to define trans-ness
           if %gt( %set_size( $tors[ $a ] ) 1 )
                if %set_and( $tors[ $a ] %set_create( %atoms(<C*>) ) )
                   setvar tors[ $a ] %set_and( $tors[ $a ] \
                        %set_create( %atoms(<C*>) ) )
                endif
                setvar tors[ $a ] %arg( 1 %set_unpack( $tors[ $a ] ) )
           for al in %range(1 %sln_atom_count( $capsln ) )
                if %eq( $tors[ $a ] %sln_rgroup_sybid( $mcore $tpat $a1 ) )
                    setvar tors[$a] $a1
                    break
                endif
           endfor
        endif
      endif
   endfor
if $CHOM!Align[DEBUG]
echo %prompt( INT 1 " " " ")
endif
endif
default $ma >$nulldev
setvar CHOM! BadRows
```

```
##
              build 3D models
##
# off we go !! Get MSS row IDS to build models for
if %streql( $1 * )
   setvar rids %table( * ROW NUM )
   setvar rids %set unpack( $1 )
endif
for rid in $rids
# get the next MSS entry to be modelled
  table examine $rid | >$nulldev
# fix NO2's (egad what a pain) because Concord & SYBYL are inconsistent
  setvar pat search2d(sln(sma)N(=0)OALL 0 y)
  while $pat
     setvar pat %sln_rgroup_sybid( $ma %arg( 1 $pat ) 1 3 )
modify bond type %bonds( %cat( %arg( 1 $pat ) "=" \
                 %arg( 2 $pat ) ) ) 2 >$nulldev
     modify atom type %arg( 2 $pat ) 0.2
     setvar pat %search2d( %sln( $ma ) N(=0)O ALL 0 y )
  endwhile
  if $CHOM!Align(DEBUG)
    label id *
  endif
# basic optimization
  switch $4
case CONCORD)
    CONCORD MOL $ma >$nulldev
# if Concord failed, we may still be awfully flat
# minimize if there are heavy atoms not part of a single aromatic system ..
    setvar noth %atoms( *-<H>>)
    setvar al %arg( 1 $noth )
    if %set_diff( "%set_create( $noth )"
        "\set_create(\frac{\pi}{atoms(\frac{\pi}{cat(\pi \{aromatic(\pi \pi \{a1\pi \pi)\}\pi \))\pi \)}")}"))")
      setvar zs %extent_3d( %cat( $ma "(*)" )
      setvar zs math( \frac{\pi}{4} arg( 5 \$zs ) - \alpha ( 6 \$zs ) )
      if %eq( $zs 0.0 )
        %unflatten( %cat( $ma "(*)" ) )
        MAXIMIN $ma DONE INTERACTIVE
      endif
    endif
case MINIMIZE)
    MAXIMIN $ma DONE INTERACTIVE >$nulldev
  endswitch
# done, if only 3d coord, but for topomeric CoMFA ..
  if %streql( %substr( $3 1 1 ) "A" )
# find any arbitrary 2D hit
    setvar pat %search2d( %cat( %sln( $ma ) ) $capsln NoDup 0 y )
    if %not( $pat )
        setvar CHOM!BadRows %set_or( "$CHOM!BadRows" $rid )
        echo $capsin not found in molecule for Row $rid .. skipping
        goto next1
```

```
endif
   setvar pat %arg(1 $pat )
   setvar allpatats %set_create( %sln_rgroup_sybid( $ma $pat \
        %range( 1 %sln_atom_count( $capsln ) ) )
# collect all appropriate heavy atoms for FIT and torsions
    setvar mat1
    setvar mat2
    setvar schns
    for a in $hvinpat
        setvar mat1 $mat1 $patats[ $a ]
        setvar sybat %sln_rgroup_sybid( $ma $pat $a )
        setvar mat2 $mat2 $sybat
# are there heavy atom neighbors to FIT also (and generate torsion lists)?
        if $patats($a)(YS)
           setvar ans %set_diff( %set_create( \
                %atom_info( $sybat NEIGHBORS ) ) $allpatats )
           setvar ans %atoms($ans-<H>)
           setvar i 1
           for p in %set_unpack( $patats[$a][YS] )
# add heavy atom neighbors to FIT list
              if %arg( $i $ans )
                setvar mat1 $mat1 $p
                setvar mat2 $mat2 %arg( $i $ans )
# generate another torsion for CHOM!alltrans
                setvar schns $schns %cat( $sybat "," \
%sln rgroup sybid( $ma $pat $tors[ $a ] ) "," %arg( $i $ans ) )
              endif
              setvar i %math( $i + 1 )
           endfor
        endif
    endfor
    setvar dofit MATCH cat( mcore "(" set_create( mat1 ) ")" ) \
        %cat( $ma "(" %set_create( $mat2 ) ")" )
    $dofit >$nulldev
if $CHOM!Align[DEBUG]
  echo %prompt( INT 1 " " " ")
endif
# do FIT
    if %gt( $MATCH_RMS $CHOM!Align[ FITRMS ] )
        setvar CHOM!BadRows %set or ( "$CHOM!BadRows" $rid )
        echo Bad geometric alignment (MATCH_RMS = $MATCH_RMS) for Row $rid .. sk
        goto next1
    endif
# side chain alignments ..
    switch $CHOM!Align[ ALICYC ]
case User Macro)
       $CHOM!Align[ ALIDATA ] $ma $CHOM!ALIGN[ MCORE ]
case All_trans)
case With_Templates)
        setvar nojrings TRUE
        setvar rbds %set_create( %bonds({rings()}) )
       for i in $schns
           setvar jbds %set unpack( $i )
# can set "side chain" bonds only if connecting bond is not cyclic
           if %set_and( "$rbds" "%bonds( %cat( %arg( 3 $jbds ) = \
```

```
%arg( 1 $jbds ) ) " )
                 setvar nojrings
                 CHOM!AllTrans $jbds
           endif
        endfor
if $CHOM!Align(DEBUG)
  echo %prompt( INT 1 " " " )
endif
        if %streql( $CHOM!Align[ ALICYC ] With Templates )
           setvar f %open( $CHOM!Align[ ALIDATA ] "r" )
           setvar buff %read( $f )
           setvar slnma %cat( %sln( $ma ) )
           while $buff
# each line of text should have pattern, SLN IDs for the 4 torsion atoms,
        and a torsion value to set
              if %eq( %count( $buff ) 5 )
                 setvar torpat %search2d( $slnma %arg( 1 $buff ) NoDup 0 y )
                 for t in $torpat
        MODIFY TORSION %sln_rgroup_sybid( $ma $t %arg( 2 $buff ) \ %arg( 3 $buff ) %arg( 4 $buff ) ) %arg( 5 $buff ) >$nulldev
                 endfor
             endif
           endwhile
           %close( $f )
        endif
: :
    endswitch
  endif
# do a bump check?
  if $CHOM!Align[BUMPS]
     if %atoms({bumps(*,*)})
        setvar CHOM!BadRows %set_or( "$CHOM!BadRows" $rid )
        echo Bad steric contacts in aligned conformer for Row $rid .. skipping
        qoto next1
     endif
  endif
# partial charges ..
  switch $CHOM!Align[ CHARGE ]
case None)
case User Macro)
    exec $CHOM!Align[ CHARGEDATA ] $ma
case )
    CHARGE $ma COMPUTE $CHOM!Align[ CHARGE ] | >$nulldev
 endswitch
# put conformer away
  switch %substr( $2 1 3 )
case SYB)
    database add $ma r >$nulldev
case )
    %wcell( $rid $newc %cat( %cat( %sln( $ma FULL CHARGE ) ) ) ) >$nulldev
 endswitch
```

```
echo Built row $rid
next1:
endfor
if %streql( %substr( $3 1 1 ) "A" )
   copy $mcsav $mcore
   zap $mcsav
endif
if $CHOM!Align[BUMPS]
   TAILOR SET GENERAL bumps_contact_distance $tailor bumps save | |
# done, restore initial EXAMINE settings
set CGQ TIMEOUT $cgq save
setvar TAILOR!EXAMINE!AFFECT_SUBSET
                                            $AFFECT_SUBSET_save
setvar TAILOR!EXAMINE!EXAMINE TAILOR MODE
                                           $EXAMINE_TAILOR MODE save
setvar TAILOR!EXAMINE!HIGHLIGHT MSS
                                            $HIGHLIGHT MSS save
setvar TAILOR!EXAMINE!INFORM
                                           $INFORM save
setvar TAILOR!EXAMINE!INPUT MODE
                                           $INPUT MODE save
setvar TAILOR!EXAMINE!RELATE
                                           $RELATE save
setvar TAILOR!EXAMINE!SHOW_MOLECULE
                                           $SHOW_MOLECULE_save
MAXIMUM_ITERATIONS %arg( 2 $max_save ) | |
# update row and column information
if %streql( %substr( $2 1 3 ) NEW )
# make any new conformer column become the source of molecules
   TABLE CONF %table( * COL COUNT )
   CHOM!UPDATE_ROW_SEL $CHOM!CID_Last
   setvar CHOM!CID_Last %math( $CHOM!CID_Last + 1 )
  CHOM! UPDATE_ROW_SEL
endif
# Section I-B. Generates the topomeric conformation of the 3D model
@macro ALLTRANS chom
# assumes default molecule, takes argument atoms $1 and $2
# where $1 is the JOINed atom of the core, $2 is the atom that
   the rest of the substituent is to be trans to,
  and $3 is the JOINed atom of the substituent
# starts from that atom and sets all side chains
# to a topomeric conformation
localvar bds b bdset a1 a2 tmp sbonds sats rbond pbds torsion ringbonds doit
# check input for legality
  setvar tmp %set_create( %atom_info( $1 NEIGHBORS ) )
  echo Bad input to ALLTRANS (atoms $2 $3 not bonded to $1)
    return
```

```
endif
# save key bonds
   setvar rbond %bonds( %cat( $3 "=" $1 ) )
   setvar sats %conn_atoms( $3 $1 )
   if %not($sats)
      echo No substituent atoms found in ALLTRANS
   endif
   setvar sats $3 $sats
   setvar sbonds %set_create( %bonds( \
        %cat( "{TO_ATOMS(" %set_create($sats) ")}" )) )
# define the other bonds that might need adjusting
   setvar bds %set_create( %bonds( (*-{RINGS()})&<1> ) )
   setvar bds %set_and( "$sbonds" "$bds" )
   if %not($bds)
      return
   endif
# discard bonds to primary atoms
   setvar mval %set create( %atoms( \
        <H>+<0.2>+<F>+<I>+<Cl>+<Br>+<n.1>+<LP>+<Du>) )
   setvar pds %set_create( %bonds( %cat( "{TO_ATOMS(" $mval ")}" ) ) )
   setvar bds %set_diff( $bds $pds )
   setvar ringbonds %set_create(%bonds({RINGS()}) )
# walk all the important bonds
 for b in %set unpack( $bds )
    setvar doit TRUE
# if this is the JOIN bond, already have some info
    if %eq( $b $rbond )
     setvar a0 $2
     setvar al $1
     setvar a2 $3
# still need to be SURE we're not monovalent
     if %or( "%eq( 1 %count( %atom_info( $a1 NEIGHBORS ) ) ) " \
         "%eq( 1 %count( %atom_info( $a2 NEIGHBORS ) ) ) " )
        setvar doit
     endif
     setvar bdat %bond_info( $b ORIGIN TARGET )
setvar al %arg( 1 $bdat )
     setvar a2 %arg( 2 $bdat )
     if %or( "%eq( 1 %count( %atom_info( $a1 NEIGHBORS ) ) ) " \
         "%eq( 1 %count( %atom_info( $a2 NEIGHBORS ) ) )" )
        setvar doit
     endif
     if $doit
# which end leads to root atom? if necessary flip a1,a2 to make that one be a1
      if %set_and( "%set_create( %conn_atoms( $a2 $a1 ) )" $1 )
        setvar tmp $a1
        setvar al $a2
        setvar a2 $tmp
      endif
      setvar a0 %trans_path( $a1 $a2 $1 )
    endif
    endif
```

if \$doit

setvar a3 %trans_path(\$a2 \$a1)

```
switch %count( %set_unpack( "%set_and( "$ringbonds" \
       %set create( %bonds( %cat( $a0 "=" $a1 "," $a2 "=" $a3 ) ) ) ) " ) )
case 0)
       setvar torsion 180
case 1)
       setvar torsion 90
case 2)
       setvar torsion 60
;;
    endswitch
    modify torsion $a0 $a1 $a2 $a3 $torsion >$nulldev
endfor
/* Beginning of section I-C, C code implementing the trans_path expression gener
/*E+:SYB MGEN CONN BEST*/
                       **************
* int SYB_MGEN_CONN_BEST( identifier, nargs, args, writer )
       Dick Cramer, Apr. 9, 1995 (written for SELECTOR use)
  Expression generator that returns the atoms attached to a given
       atom, excepting the second, in a prioritized order.
  If there are two arguments, the ordering is by decreasing branch
       "size", where "size" is first any path with rings encountered, then
  number of attached atoms, then MW (paths in cycles end when an atom
* in another path is encountered.)
     If three arguments, the atom that is returned is the one that
  begins the shortest path containing the atom referred to by the
  third argument. If multiple such paths, ordering is same as for
  two arguments.
     Further prioritization of paths is by molecular weight,
     and then by lowest X, Y, Z values.

If last argument is DEBUG, all paths are written to stdout.
  User interface:
    %trans_path( a1 a2 ( a3 ) (DEBUG) )
****************
int SYB_MGEN_CONN_BEST( identifier, nargs, args, Writer )
/* following arguments contain the text supplied to the %trans_path()
 expression generator, and provide an avenue for producing text output. */
char
       *identifier;
int
       nargs;
char
       *args[];
PFI
       Writer;
# define MAX_NP 8
       struct pathrec {
         int root, nrings, chosen, nats;
         float mw, xyz[3];
         set_ptr path;
       } ;
       struct pathrec p[MAX_NP];
```

```
int retval, i, np, toroot, a1, a2, a4, a, pnow, pdone, growing,
            final_pos, area_num, new_rings, nats, nuats, elem, ncycles,
           best, debug, ringclosed;
        List Ptr
                     atom_exp_list=NIL,SYB_EXPR_ANALYZE();
        mol ptr
                     m1, m2, SYB_AREA_GET_MOLECULE();
        atom_ptr
                     arec, SYB_ATOM_FIND_REC();
/* A set_ptr data structure is a Boolean set, first word containing
its cardinality. */
                    atom_set1=NIL, a2chk = NIL, nuls = NIL, cnats = NIL,
        set_ptr
                nxcn = NIL, end_atoms = NIL, scratch = NIL,
                 SYB_ATOM_FIND_SET(), UTL_SET_CREATE();
        char
                   tempString[256];
        float
                   get_path_mw(), diff;
        void
                   get_path_xyz();
        retval = 0;
        /* Check the number of arguments */
        if ( nargs < 2 | nargs > 4 )
                UIMS2 WRITE ERROR(
                   "Error: %trans_path requires 2 to 4 arguments\n" );
                return 0;
        }
        np = 0;
        debug = (!UTL_STR_CMP_NOCASE( args[ nargs - 1], "DEBUG" ));
        toroot = (debug && nargs == 4) | (!debug && nargs == 3);
/* PARSE THE INPUT */
/* get first atom */
    if (!(atom_exp_list = SYB_EXPR_ANALYZE( SYB_EXPR_GET_ATOM_TOKEN, args[0],
        &final_pos, &area_num )))
       goto error;
    if (!(m1 = SYB_AREA_GET_MOLECULE (area num)))
       goto cleanup;
    if (!(atom_set1 = SYB_ATOM_FIND_SET ( m1, atom_exp_list)))
        goto error;
    if( atom_exp_list)
          SYB_EXPR_DELETE_RPN_LIST( atom_exp list);
    atom_exp_list = (List_Ptr) NIL;
    if(!(1 == UTL_SET_CARDINALITY(atom_set1))) {
                UIMS2 WRITE ERROR (
                  "Error: First argument must be only one atom\n");
                goto error;
   if (!(arec = SYB_ATOM_FIND_REC (m1, UTL_SET_NEXT (atom_set1, -1)) )) goto er
   a1 = arec->recno;
   UTL SET DESTROY( atom_set1 );
   atom_set1 = NIL;
/* get 2\overline{n}d atom */
   if (!(atom_exp_list = SYB_EXPR_ANALYZE( SYB_EXPR_GET_ATOM_TOKEN, args[1],
       &final_pos, &area_num ).))
      goto error;
   if (!(m2 = SYB_AREA_GET_MOLECULE (area_num)))
      goto cleanup;
   if (!(end_atoms = SYB_ATOM_FIND_SET ( m2, atom_exp_list)))
       goto error;
```

```
if( atom exp_list)
           SYB_EXPR_DELETE_RPN_LIST( atom_exp_list);
    atom exp list = (List Ptr) NIL;
    if (m1 != m2 ) {
                 UIMS2 WRITE ERROR(
                   "Error: atoms must be in the same molecule\n");
                 goto error;
    if(!(1 == UTL_SET_CARDINALITY(end_atoms))) {
                 UIMS2_WRITE_ERROR(
                   "Error: Second argument must be only one atom\n");
                 goto error;
    if (!(arec = SYB_ATOM_FIND_REC (m1, UTL_SET_NEXT (end_atoms, -1)) )) goto er
    a2 = arec->recno;
/* get 3rd atom */
 if (toroot) {
    if (!(atom_exp_list = SYB_EXPR_ANALYZE( SYB_EXPR_GET_ATOM_TOKEN, args[2],
        &final_pos, &area_num )))
       goto error;
    if (!(m2 = SYB_AREA_GET_MOLECULE (area_num)))
       goto cleanup;
    if (!(atom_set1 = SYB_ATOM_FIND_SET ( m2, atom_exp_list)))
        goto error;
    if( atom_exp_list)
          SYB_EXPR_DELETE_RPN_LIST( atom_exp_list);
    atom_exp_list = (List Ptr) NIL;
    if (m1 != m2 ) {
                 UIMS2_WRITE_ERROR(
                   "Error: atoms must be in the same molecule\n");
                goto error;
    if(!(1 == UTL SET CARDINALITY(atom_set1))) {
                UIMS2 WRITE ERROR(
                   "Error: Second argument must be only one atom\n");
                goto error;
    if (!(arec = SYB_ATOM_FIND_REC (m1, UTL_SET_NEXT (atom_set1, -1)) )) goto er
    a4 = arec->recno;
    UTL_SET_DESTROY( atom_set1 );
    atom set1 = NIL;
/* GENERATE the paths */
/* set up paths */
   if (!(a2chk = UTL_SET_CREATE( m1->max_atoms + 1 ) )) goto error;
   if (!(nuls = UTL_SET_CREATE( ml->max_atoms + 1 ) )) goto error;
if (!(cnats = UTL_SET_CREATE( ml->max_atoms + 1 ) )) goto error;
   if (!(nxcn = UTL_SET_CREATE( m1->max_atoms + 1 ) )) goto error;
   if (!(scratch = UTL_SET_CREATE( m1->max_atoms + 1 ) )) goto error;
   if (!syb_mgen_conn_att_atoms( a2chk, m1, a1 )) goto error;
   if (!UTL_SET_MEMBER( a2chk, a2 )) {
```

```
UIMS2 WRITE ERROR (
           "Error: second argument atom is not bonded to first argument atom/\n")
        goto error;
    UTL_SET_DELETE( a2chk, a2 );
    a = -1;
    np = 0;
    while (np < MAX_NP && (a = UTL_SET_NEXT( a2chk, a)) >= 0 )
        if (!(p[np].path = UTL_SET_CREATE( ml->max_atoms + 1 ) )) goto error;
        p[np].root = a;
        p(np).nrings = 0;
        UTL SET INSERT ( p[np].path, a );
        np++;
/* grow the paths */
    growing = TRUE;
    nats = 0;
    ncycles = 0;
    while (growing ) {
      nuats = 0;
      ringclosed = FALSE;
      for (pnow = 0; pnow < np; pnow++) {
        UTL_SET_COPY_INPLACE( cnats, p[pnow].path );
UTL_SET_CLEAR( nxcn );
        elem = -1;
/* accumnulate this generation of attached atoms into nxcn */
        while ( (elem = UTL_SET_NEXT( cnats, elem)) >= 0 ) {
           UTL_SET_CLEAR( nuls );
            if (!syb_mgen_conn_att_atoms( nuls, ml, elem )) return( FALSE );
           UTL_SET_DELETE( nuls, al );
UTL_SET_DIFF_INPLACE( nuls, end_atoms, nuls );
           UTL SET OR INPLACE( nxcn, nuls, nxcn );
           UTL SET_DIFF_INPLACE( nxcn, p[pnow].path, nxcn );
        UTL_SET_OR_INPLACE( p[pnow].path, nxcn, p[pnow].path );
/* remove and mark ring closures when growing out */
        if (!toroot) for (pdone = 0; pdone < np; pdone++ ) if (pdone != pnow) {</pre>
           UTL_SET_AND_INPLACE( p[pnow].path, p[pdone].path, a2chk );
           if ((new_rings = UTL_SET_CARDINALITY( a2chk ))) {
/* we have ring closure(s) */
                 p[pnow].nrings += new_rings;
                 p[pdone].nrings += new_rings;
                 ringclosed = TRUE;
UTL_SET_OR_INPLACE( end_atoms, a2chk, end_atoms );
/* if pdone < pnow, two branches are now same lengths, drop common atom from bot</pre>
        but if >, branches are different, and must avoid repeated closing */
                 if (pdone < pnow) {
   /* remove atom(s) in the previous branch because paths are really same length
                    UTL_SET_DIFF_INPLACE( p[pdone].path, a2chk, p[pdone].path );
                    UTL_SET_DIFF_INPLACE( p[pnow].path, a2chk, p[pnow].path );
                 else {
/\star must identify and mark each atom in nxcn that is attached to a2chk atom \star/
                    elem = -1;
                    while ( (elem = UTL_SET_NEXT( a2chk, elem)) >= 0 ) {
                         UTL_SET_CLEAR( scratch );
                         if (!syb_mgen_conn_att_atoms( scratch, m1, elem ))
                                  return( FALSE );
                         UTL_SET_AND_INPLACE( scratch, nxcn, scratch );
```

```
UTL SET OR INPLACE( end atoms, scratch, end_atoms );
                    }
                }
           }
        }
/* done growing paths if no more atoms added to any path .. */
      for (pdone = 0, nuats = 0; pdone < np; pdone++ )
     nuats += UTL_SET_CARDINALITY( p[pdone].path );</pre>
      if (nuats<=nats && !ringclosed) growing = FALSE;
      nats = nuats;
  .. or looking for the 4th atom and found it .. \star/
      if (toroot) for (pdone = 0; pdone < np; pdone++ )
          if (UTL SET_MEMBER( p[pdone].path, a4 )) growing = FALSE;
  .. or after 100 atom layers out regardless */
      ncycles++;
      if (ncycles >= 100) growing = FALSE;
/* debugging */
   if (debug) for (pdone = 0; pdone < np; pdone++) {
    sprintf( tempString, "Path %d (%d rings, from %d): ",</pre>
                 pdone+1, p[pdone].nrings, p[pdone].root );
        UBS_OUTPUT_MESSAGE( stdout, tempString );
        ashow(p[pdone].path, m1);
/* compute the path properties */
   for (pdone = 0; pdone < np; pdone++) {
  /* mark as already chosen any path that can't be an answer */
        p[pdone].chosen = toroot && !UTL_SET_MEMBER(p[pdone].path, a4);
        p[pdone].nats = UTL_SET_CARDINALITY( p[pdone].path );
        p(pdone).nrings = p[pdone].nrings ? 1 : 0;
        p[pdone].mw = 0.0;
        p[pdone].xyz[0] = p[pdone].xyz[1] = p[pdone].xyz[2] = 0.0;
/* return the best result */
   best = 0;
   for (pdone = 1; pdone < np; pdone++) {</pre>
        if (toroot) {
           if (p[best].chosen && !p[pdone].chosen) best = pdone;
/* looking backward along chain, always grow away from more negative coord value
           if (!p[best].chosen && !p[pdone].chosen) {
                 get_path_xyz( p[pdone].root, m1, p[pdone].xyz );
                 get_path_xyz( p[best].root, m1, p[best].xyz );
                 for (i = 0; i < 3; i++) {
                    diff = p[pdone].xyz[i] - p[best].xyz[i];
                    if (diff < -0.1)
                         best = pdone;
                         break;
                    if (diff > 0.1 ) break;
/* checking other coords if basically tied at this coord */
        else
          if (p[pdone].nrings && !p[best].nrings) best = pdone;
          else if (p[pdone].nats > p[best].nats) best = pdone;
          else if (p[pdone].nats == p[best].nats) {
           p[pdone].mw = get_path_mw( p[pdone].path, m1, p[pdone].mw );
           p[best].mw = get_path_mw( p[best].path, ml, p[best].mw );
```

```
if (p[pdone].mw > p[best].mw) best = pdone;
        }
   arec = SYB ATOM FIND REC( ml, p[best].root );
   sprintf(tempString,"%d", arec->id );
   if(!(*Writer)(tempString)) goto error;
   retval = TRUE;
error:
cleanup:
    if( atom_exp_list)
           SYB_EXPR_DELETE_RPN_LIST( atom_exp_list);
    if (atom set1)
            UTL SET DESTROY (atom set1);
    if(end_atoms)
            UTL_SET_DESTROY(end_atoms);
    if (a2chk)
            UTL SET DESTROY (a2chk);
    if (nuls)
            UTL_SET_DESTROY(nuls);
    if (nxcn)
            UTL SET DESTROY(nxcn);
    if (cnats)
           UTL_SET_DESTROY(cnats);
    if(scratch)
           UTL_SET_DESTROY(scratch);
    return( retval );
}
static int syb_mgen_conn_att_atoms( aset, m, atid )
/* ors atoms attached to atm into aset */
/* WORKS STRUCTLY WITH RECNOS */
set ptr aset;
mol_ptr m;
int atid;
   atom ptr at, SYB ATOM FIND ID();
   List_Ptr tohs, UTL_LIST RETRIEVE P();
   atom_ptr toh, SYB_ATOM_FIND_REC();
   acon ptr conn1;
   int nbytes1;
   at = SYB ATOM FIND REC( m, atid );
   tohs = at->conn_atom;
   while (tohs) {
        tohs = UTL_LIST_RETRIEVE_P( tohs, &conn1, &nbytes1);
toh = SYB_ATOM_FIND_REC( m, conn1->target );
        UTL_SET_INSERT( aset, toh->recno );
   return( TRUE );
static float get path mw( aset, m, mw )
/st returns the total atomic weight of all atoms in aset st/
set_ptr aset;
mol ptr m;
float mw;
```

```
int elem = -1;
  float ans = 0.0;
  atom_ptr at, SYB_ATOM_FIND_REC();
  fpt SYB ATAB ATOMIC WEIGHT();
  if (mw) return( mw );
  elem = -1;
  while ( (elem = UTL_SET_NEXT( aset, elem)) >= 0 ) {
     at = SYB_ATOM_FIND_REC( m, elem );
     ans += (float) SYB_ATAB_ATOMIC_WEIGHT( at->type );
  return( ans );
}
static void get_path_xyz( aid, m, mw )
/* returns the xyz of the supplied atom */
int aid;
mol_ptr m;
float mw[3];
  int i;
  atom_ptr at, SYB_ATOM_FIND REC();
  if (mw[0]) return;
  at = SYB_ATOM_FIND_REC( m, aid );
 for (i = 0; i < 3; i++) mw[i] = at->xyz[i];
  return;
static int ashow( aset, m )
/* for interactive debugging, shows a set's membership in terms of atom ID */
set_ptr aset;
mol_ptr m;
     char buff[1000], *b;
     atom ptr at, SYB ATOM FIND REC();
     int elem;
     *buff = '/0';
     b = buff;
     elem = -1;
     while ( (elem = UTL_SET_NEXT( aset, elem)) >= 0 ) {
           at = SYB_ATOM_FIND_REC( m, elem );
           sprintf( b, " %d", at->id );
           b = buff + strlen( buff );
     sprintf( b, "\n" );
     UBS_OUTPUT_MESSAGE( stdout, buff );
/* BEGINNING OF SUBROUTINES I-D. Calculation of attenuated fields */
/*+E:QSAR_FIELD EVAL RB ATTEN()*/
/* int QSAR_FIELD_EVAL_RB_ATTEN( molp, stfldp, elfldp, regp, no_st, no_el, ctp )
   Dick Cramer
                   May 13, 1995
```

```
"Standard CoMFA" -- except that the contribution of any atom
      to the field falls off with an inverse power of its distance
      from a root atom, measured in NUMBER OF ROTATABLE BONDS!
         This means also that each individual atom's contribution
     has a similarly scaled upper bound, rather than checking
     the upper bound only for the sum over all atoms.
/* This procedure computes vdW 6-12 steric values at each point in region
/* and the electrostatic interactions (initially assuming 1/r dielectric).
    NOTE:: initially ignoring space averaging, other user knobs.
    note:: assuming valid input here; error checking higher up !
   Input:
/*
              - molecule pointer, molecule to place in region.
     molp
             - steric field pointer, where values will be placed.
     stfldp
     elfldp - electrostatic field pointer, where values will be placed.
              - region pointer, locations where values are to be evaluated.
     regp
              - flag to skip steric evaluations
     no st
              - flag to skip electrostatic evaluations

    ComfaTopPtr, for dummy/lp values

/* Returns 0 on failure, 1 otherwise.
/******************
/*+E:QSAR FIELD EVAL RB ATTEN()*/
int QSAR_FIELD_EVAL_RB_ATTEN ( molp, stfldp, elfldp, regp , no st, no el, ctp)
mol_ptr molp;
FieldPtr stfldp, elfldp;
RegionPtr regp;
int no_st, no_el ;
ComfaTopPtr ctp;
BoxPtr box;
atom_ptr at, SYB_ATOM_FIND_ID();
int pid, b, ix, iy, iz, nat, vol_avg, repulsive;
fpt *steric, *elect, SYB_ATAB_VDW_RADII();
fpt diff, dis, dis2, x, y, z, sum_steric, sum_elect;
fpt dis6, dis12, repuls_val, offs[9][3], atm_ste, atm_ele;
fpt *charge, *ctemp, *coord, *ftemp, *wt, scale_vol_avg, atm_steric, atm_elect;
int *atyp, *itemp, dohbd, dohba, ishbd, retval, dielectric, off, atid;
static fpt hbond_scal;
fpt hbond_A, hbond_B, *AtWts = NIL, *QSAR_FIELD_RB_WTS();
int *HAs, *HDs, *HAp, *HDp; /* sets would be more efficient but slower */
int do_steric, do_elect;
set_ptr hdonor, SYB_HBOND_DONORS(), pset = NIL, aset = NIL;
#define Q2KC 332.0
#define MIN SQ DISTANCE 1.0e-4
/* ^^^ any atom within 10-2 Angstroms is hereby zapped !
       this is about it: 10<sup>6</sup> / 10<sup>-24</sup> is close to overflow!
   ftemp = NIL; ctemp = NIL; itemp = NIL; retval = FALSE; HAs = NIL; HDs = NIL;
   hdonor = NIL;
/* for now, make root atom the one closest to 0,0,0 */
   for (nat = 1; nat <= molp->natoms; nat++) {
```

```
at = SYB ATOM_FIND_ID( molp, nat );
      dis2 = a\overline{t} - xy\overline{z}[0] + at - xyz[0] + at - xyz[1] + at - xyz[1] +
                at->xyz[2] * at->xyz[2];
      if (nat == 1 || dis2 < dis) {
        dis = dis2;
        atid = nat;
      }
/* following is specific to topomeric fields */
if (!(AtWts = QSAR_FIELD_RB_WTS( molp, atid ) )) goto cleanup;
if (!no el)
  {dielectric = elfldp->dielectric ;
  vol_avg = elfldp->vol_avg_type;
  scale_vol_avg = elfldp->scale_vol_avg;
  repulsive = elfldp->repulsive;
  repuls_val=repexp[repulsive]; elect = elfldp -> field_value;}
 if (!no st)
  {vol avg = stfldp->vol_avg_type;
   scale vol avg = stfldp->scale_vol_avg;
  repulsive = stfldp->repulsive;
   repuls_val=repexp(repulsive); steric = stfldp -> field_value;}
 if (!(ftemp = (fpt *) UTL_MEM_ALLOC(3*sizeof(fpt)*molp->natoms))) goto cleanup;
 if (!(ctemp = (fpt *) UTL MEM ALLOC( sizeof(fpt)*molp->natoms))) goto cleanup;
    (!(itemp = (int *) UTL_MEM_ALLOC( sizeof(int)*molp->natoms))) goto cleanup;
if (!(HAs = (int *) UTL MEM_ALLOC( sizeof(int)*molp->natoms))) goto cleanup;
if (!(HDs = (int *) UTL_MEM_ALLOC( sizeof(int)*molp->natoms))) goto cleanup;
/* get just those H's which are capable of Hbonding */
 if (!(hdonor = SYB_HBOND_DONORS( molp, NIL ) )) goto cleanup;
 for (coord=ftemp,atyp=itemp,charge=ctemp,HAp=HAs,HDp=HDs, nat=1;
                nat<=molp->natoms;nat++)
  { if (NIL == (at = SYB_ATOM_FIND_ID(molp, nat) ) ) goto cleanup;
    *coord++ = at->xyz[0];
    *coord++ = at->xyz[1];
    *coord++ = at->xyz[2];
              = at - > type - 1 ;
    *atyp++
    *charge++ = at->charge;
              = SYB ATAB HBOND_ACCEPT(at->type) ;
    ++qAH*
              = UTL_SET_MEMBER(hdonor, at->recno) ;
    *HDp++
 for (b=0; b<regp->n_boxes; b++) {
  box = & regp->box_array(b);
  dohbd = (SYB_ATAB_ATOMIC_NUMBER( box->atom_type) == 1) &&
        (box->pt_charge == 1.0);
  dohba = (SYB ATAB ATOMIC_NUMBER( box->atom_type ) == 8) &&
        (box->pt_charge == -1.0);
  if (dohbd || dohba)
        if (!TAILOR_STORE_IT_HERE( "TAILOR!FORCE_FIELD!HBOND_RAD_SCALING",
                &hbond_scal, 1)) goto cleanup;
        hbond_A = pow( hbond_scal, 6.0 );
        hbond B = hbond A * hbond A;
  if (vol_avg)
    QSAR_FIELD_EVAL_GETOFF(offs,box->stepsize,vol_avg,scale_vol_avg);
  if (!no st)
    QSAR_FĪELD_VDWTAB ( box -> atom_type, repuls_val, ctp->du_lp_steric );
  for (i\overline{z}=0, z=box->lo[2]; iz < box->nstep[2]; iz++, z += box->stepsize[2])
```

```
for (iy=0, y=box->lo[1]; iy < box->nstep[1]; iy++, y += box->stepsize[1])
   for (ix=0, x=box->lo[0]; ix < box->nstep[0]; ix++, x += box->stepsize[0])
     nat<molp->natoms;
       nat++, wt++)
      if ( ( *atyp == DUMMY-1 || *atyp == LP-1 ) && !ctp->du_lp_elect )
         *charge = 0.0; /* set charge to 0 since ignoring Du/lp */
      if (!vol_avg) /* the "normal" case */
       dis2 = x - *coord++ ;
       dis2 *= dis2;
       diff = y - *coord++ ;
       diff *= diff;
       dis2 += diff;
       diff = z - *coord++;
       diff *= diff;
       dis2 += diff;
       if (!no el && elfldp->zap el==2 && do elect)
         dis = sqrt( dis2 );
         if ( dis < SYB_ATAB_VDW_RADII( *atyp+1 ) ) {</pre>
/* no shortcircuits! */
            *elect++ = 0.0;
            do elect = FALSE;
*/
         }
       if ( dis2 < MIN_SQ_DISTANCE ) {
          if (!no_st)
             /* if atom has no steric value, we don't care about
                MIN_SQ_DISTANCE since it has no contribution anyway */
             if (vdw_a[*atyp] != 0.0 && vdw_b[*atyp] != 0.0) {
               /* set sterics to its max value at current grid pt. */
               atm_steric = (*wt) * stfldp->max_value;
          if (!no_el && do_elect)
             if ( !no_st && !do_steric && elfldp->zap_el ) {
                *elect++ = DAB_F_MISSING;
              else if ( *charge != 0.0 ) {
                if ( *charge > 0.0 )
                  atm_elect = (*wt) * elfldp->max_value;
                else atm_elect = (*wt) * -elfldp->max value;
              }
          if ( !do_elect && !do_steric )
            break;
                     /* break out of loop since neither el. or st.
                        need to be calculated for this grid point */
           /* setting dis2 to 1 (an arbitrary no.) will prevent a zero
             divide in the sum_steric or sum_elect calculations below */
           dis2 = 1.0;
       if ( ! no_st && do_steric ) {
        dis6 = d\overline{i}s2 * dis\overline{2} * dis2;
```

```
dis12= dis6 * dis6 ;
       if (repulsive)
         dis12 = (repulsive==1) ? dis12 / dis2 : dis12 / dis2 / dis2;
       if (dohbd && *HAp)
              atm steric = hbond B * vdw b[*atyp]/dis12 -
                       hbond_A * vdw_a[*atyp]/dis6;
          else if (dohba && *HDp)
              atm_steric = hbond_B * vdw_b[*atyp]/dis12 -
                       hbond_A * vdw_a[*atyp]/dis6;
          else
              atm_steric = vdw_b[*atyp]/dis12 - vdw a[*atyp]/dis6 ;
       HAp++; HDp++;
       atm_steric = atm_steric > stfldp->max_value ? stfldp->max_value
              : atm_steric;
       atm_steric *= (*wt);
      if ( ! no_el && do_elect ) {
       atm elect = *charge++ /
                       ( dielectric ? sqrt(dis2) : dis2 ) ;
       atm_elect = atm_elect > elfldp->max_value ? elfldp->max_value
              : atm elect;
       atm_elect = atm_elect < -(elfldp->max_value) ? -(elfldp->max_value)
              : atm_elect;
       atm_elect *= (*wt);
       sum_elect += atm_elect;
       atyp++;
       sum_steric += atm_steric;
     else
     for (off=0;off<9;off++)</pre>
    coord += 3;
    atyp ++
    charge ++ ;
    HAp ++
    HDp ++
      } /* atom loop */
doneatoms:
   if ( do_steric || do_elect ) {
  if (vol_avg) { sum_elect /= 9.0; sum_steric /= 9.0; }
     if ( !no el && do elect )
      { *elect = sum_elect * box-> pt_charge * Q2KC
        if ( *elect > elfldp->max_value ) *elect = elfldp->max value;
        else if ( *elect < - elfldp->max value ) *elect =
              elfldp->max_value;
          transform_field(elfldp->max_value,elect,ctp);
          elect ++;
    if ( !no st && do_steric )
      { *steric = sum_steric ;
       if ( *steric > stfldp->max_value)
         { *steric = stfldp->max_value;
           if (!no_el && elfldp->zap_el==1 ) *(elect-1) = DAB F MISSING; }
       transform_field(stfldp->max_value, steric, ctp);
       steric ++ ; }
  } /* points in box loop */
```

```
} /* boxes loop */
  retval = TRUE;
cleanup:
  if ( itemp) UTL_MEM_FREE( itemp);
  if (ftemp) UTL_MEM_FREE(ftemp);
  if ( ctemp) UTL MEM FREE ( ctemp);
  if (HAS) UTL MEM_FREE( HAS );
  if (HDs) UTL_MEM_FREE( HDs );
  if (hdonor) UTL_SET_DESTROY( hdonor );
if (AtWts) UTL_MEM_FREE( AtWts );
  if (pset) UTL_MEM_FREE( pset );
  if (aset) UTL MEM_FREE( aset );
return retval;
#undef Q2KC
#undef MIN_SQ_DISTANCE
static fpt *QSAR_FIELD_RB_WTS( molp, rootid )
/* generates rotational-bond wts for each atom */
mol_ptr molp;
int rootid;
/* pseudo code for FIELD_RB_WTS()
   while saw new atoms
     uncover atoms that stopped last shell growth
     grow next "rotational shell"
     while adding to shell
        for each atom in shell
            get neighbors not seen
            for each neighbor
               if bond is rotatable (acyclic, >1 attached atom, not =,am,#)
                 cover all other atoms attached to atom for this shell
               add it to shell
*/
   fpt *ansr = NIL, *vals = NIL, factor, nowfact = 1.0;
                 found, aggcount, atid, aggid, loop, size;
   int
                 aggats = NIL, allats = NIL, nuls = NIL, endatms = NIL, end_cands
   set_ptr
                 root, SYB_ATOM_FIND_REC(), at, atrec ;
   atom ptr
                b, SYB_BOND_FIND_REC();
   bond ptr
                toats, UTL_LIST_RETRIEVE_P();
   List Ptr
   acon_ptr
                cptr:
   char
                 tempString[200];
                 ashow(), qsar_field_attached_atoms();
   void
   if (!( vals = (fpt *) UTL MEM ALLOC( sizeof(fpt)*molp->natoms))) return( NI
   if (!UIMS2 VAR GET TOKEN( "TAILOR!COMFA!AGGREG DESCALE",
        &factor ) ) return( NIL );
   if (!(allats = UTL_SET_CREATE( molp->max_atoms + 1 ) )) goto cleanup;
if (!(aggats = UTL_SET_CREATE( molp->max_atoms + 1 ) )) goto cleanup;
   if (!(nuls = UTL_SET_CREATE( molp->max_atoms + 1 ) )) goto cleanup;
   if (!(endatms = UTL_SET_CREATE( molp->max_atoms + 1 ) )) goto cleanup;
   if (!(end_cands = UTL_SET_CREATE( molp->max_atoms + 1 ) )) goto cleanup;
   if (!( root = SYB_ATOM_FIND_REC( molp, rootid ) )) goto cleanup;
   UTL_SET_INSERT( aggats, root->recno );
UTL_SET_INSERT( allats, root-> recno );
   aggcount = loop = 1;
```

```
while (TRUE) {
        while (TRUE) {
           aggid = -1;
           while ((aggid = UTL_SET_NEXT( allats, aggid )) >= 0 ) {
                 UTL SET CLEAR ( nuls );
                 qsar_field_attached_atoms( nuls, molp, aggid );
UTL_SET_DIFF_INPLACE( nuls, allats, nuls );
UTL_SET_DIFF_INPLACE( nuls, endatms, nuls );
/* identifying any atoms that terminate this aggregate */
                 atid = -1;
                 while ((atid = UTL_SET_NEXT( nuls, atid )) >= 0 ) {
                   if (!( at = SYB_ATOM_FIND_REC( molp, atid ) )) goto cleanup;
/* skipping monovalent atoms */
                   if (at->nbond > 1) {
/* find bond record that attaches to aggid */
                      toats = at->conn_atom;
                      found = FALSE;
                      while (toats && !found ) {
                          toats = UTL_LIST_RETRIEVE_P( toats, &cptr, &size );
                          found = (cptr-> target == aggid );
                      if (!found) goto cleanup;
                      b = SYB BOND_FIND_REC (molp, cptr->bond_rec);
                      if ( !(b->status & BOND_V_IRING) && !(b->status & BOND_V_ERI
                                   && (b->type == SYB_BTAB_MNEM_TO_TYPE("1") ) ) {
/* have an end-of-aggregate atom, mark as end atoms all other attached atoms */
                          UTL_SET_CLEAR( end_cands );
                          qsar_field_attached_atoms( end_cands, molp, at->recno );
                          UTL SET DELETE ( end cands, aggid );
                          UTL_SET_OR_INPLACE( endatms, end_cands, endatms );
                 UTL SET OR INPLACE ( aggats, nuls, aggats );
            if (UTL_SET_CARDINALITY( aggats ) <= aggcount ) break;</pre>
            aggcount = UTL SET CARDINALITY ( aggats );
            UTL_SET_OR_INPLACE( allats, aggats, allats);
/* debugging stuff .. */
         sprintf( tempString, "Aggregate %d (weight = %f ):", loop, nowfact );
        UBS OUTPUT MESSAGE( stdout, tempString );
        ashow( aggats, molp );
/* if no atoms added, we are done! */
         if (UTL SET EMPTY( aggats )) break;
/* record scaling factor for atoms in this aggregate */
        atid = -1;
        while ((atid = UTL_SET_NEXT( aggats, atid )) >= 0 ) {
   if (!(atrec = SYB_ATOM_FIND_REC( molp, atid ))) goto cleanup;
             vals[ (atrec->id)-1 ] = nowfact;
         UTL_SET_OR_INPLACE( allats, aggats, allats );
         UTL_SET_CLEAR( aggats );
         UTL SET CLEAR ( endatms );
         aggcount = 0;
         nowfact *= factor;
         loop++;
   }
```

```
ansr = vals;
cleanup:
   if (aggats) UTL_SET_DESTROY( aggats );
   if (allats) UTL_SET_DESTROY( allats );
   if (endatms) UTL SET DESTROY( endatms );
   if (end_cands) UTL_SET_DESTROY( end_cands );
   if (nuls) UTL_SET_DESTROY( nuls );
   return(ansr);
static void qsar_field_attached_atoms( aset, m, atid )
/* ors atoms attached to atm into aset */
/* WORKS STRUCTLY WITH RECNOS */
set_ptr aset;
mol_ptr m;
int atid;
   atom ptr at, SYB_ATOM_FIND_ID();
   List_Ptr tohs, UTL_LIST_RETRIEVE_P();
   atom ptr toh, SYB ATOM FIND REC();
   acon_ptr conn1;
   int nbytes1;
   at = SYB ATOM_FIND_REC( m, atid );
   tohs = at->conn_atom;
   while (tohs) {
        tohs = UTL_LIST_RETRIEVE_P( tohs, &conn1, &nbytes1);
        toh = SYB ATOM FIND REC( m, conn1->target );
        UTL SET INSERT( aset, toh->recno );
   return;
}
static void ashow( aset, m )
/* for interactive debugging, shows a set's membership in terms of atom ID */
set ptr aset;
mol ptr m;
     char buff[1000], *b;
     atom_ptr at, SYB_ATOM_FIND_REC();
     int elem;
     *buff = '/0';
     b = buff;
     elem = -1;
     while ( (elem = UTL SET NEXT( aset, elem)) >= 0 ) {
           at = SYB_ATOM_FIND_REC( m, elem );
           sprintf( b, " %d", at->id );
b = buff + strlen( buff );
     sprintf(b, "\n");
     UBS_OUTPUT_MESSAGE( stdout, buff );
}
```

```
Section II-A. SPL invoked shell for computing the diagonal defining the
#
        "best" triangle, e.g., the one with the highest density of points below.
@expression_generator LRT_FAST
# Usage:
   lrt_fast rows descriptor_cols bio_col [pls flags like scaling in quotes]
        rows (*) - rows to take
        descriptor_cols - which columns are the neighborhood metrics
        bio_col - which column has the bio (probably log bio) data
        [...] - if need to SCAL NONE or anything like that, do it here
 returns a line of the form
     3.09691 / 0.000546509 = 5666.71 - 496 : 496 :: 15.6981 : 15.6989
        max bio difference
                   optimal distance division for max bio
                                slope
#
#
                                         ^number in the lrt
                                               ^total number
                                                     `area in the lrt
                                                                  `total area
# Significance is related to whether ratio of numbers is
    much above ratio of areas.
 globalvar SAMPLS_IN_PROGRESS DONE_CHECKED_OUT
 localvar hold distname rows cols bio
 setvar rows %promptif("$1" ROW_EXP "*" "Rows to use in lrt")
 setvar cols %promptif("$2" COL_EXP "COMFA*" "Columns of mol descriptors")
 setvar bio %promptif("$3" COL_EXP "LOGBIO" "Column of bio data")
 setvar hold SAMPLS IN PROGRESS
 setvar SAMPLS_IN_PROGRESS $bio
 setvar distname TAILOR!HIER!DIST FNAME
 setvar TAILOR!HIER!DIST FNAME 1rt fort.3
# here the information is computed and written to a file
        whose name is passed in via a TAILOR value
 QSAR ANA DO I >$NULLDEV
                          $rows $cols HIER $4 |
setvar SAMPLS IN PROGRESS $hold
setvar TAILOR!HIER!DIST_FNAME $distname
# contents of the file are returned to the caller
 setvar hold %system("cat lrt_fort.3")
 %return( "$hold" )
# Section II-B. SPL script for computing the significance of the distribution
       found by 1rt fast
@expression generator dochi
# computes the chi-square statistic for the number of points below
# the diagonal, null hyptheses being the area fraction of the total.
       To be called as: %dochi( %lrt_fast( ) ), i.e., its inputs
#
```

```
# are exactly the output of %1rt fast as described in the 1rt_fast header.
   setvar expected %math( $9 * $11 / $13 )
   setvar sq %math( $7 - $expected )
   setvar sq %math( $sq * $sq / $expected )
   %return( $sq )
/* Section II-C. Computes the best diagonal in the "virtual graph" of biological
distances vs property differences. */
int QSHELL_HIER_LRT(table, biocol, dmat, nrow, order, lmsg)
char *table;
int biocol, /* column in MSS with biological data */
    nrow, /* dimension of dmat and order */
    *order; /* array of row IDs to consider */
fpt *dmat; /* distance matrix for property distances */
char *lmsg; /* file name for results */
fpt *p, *q, fabs(), bmax;
int i,j, count, status_array;
char *fpt_colname;
FILE *out, *UTL FILE FOPEN();
  /* need to get the bio values
     In the n^2 we can repack into n(n-1)/2 then add the n bio values
     and finish with the bio distances */
     No error handling. Better be data in those rows!
  */
 for (count=0, i=0; i<nrow; i++)
  for (j=0; j<i; j++)
    dmat(count++) = dmat(i*nrow + j);
q = p = dmat + ( (nrow-1) * nrow) / 2;
TBL_ACCESS_INDEX_TO_COLNAME(table, biocol-1, &fpt_colname);
TBL_GRAB_INIT_FPTS(table, 1, &fpt_colname);
 for (i=0;i<nrow;i++,p++)
   TBL_GRAB_GET_FPTS_INV(order[i]-1, &status_array, p);
TBL_GRAB_COMPLETE_FPTS();
bmax = 0.0;
for (count=0, i=0; i<nrow; i++)
 for (j=0; j<i; j++, count++)
    if (p[count] = fabs(q[i] - q[j])) > bmax) bmax = p[count];
out = UTL_FILE_FOPEN(lmsg, "w");
QSHELL_HIER_DO_LRT(out, count, dmat, p, bmax);
UTL FILE FCLOSE(out);
```

```
int QSHELL_HIER_DO_LRT( out, index, xsort, ysort, bmax )
FILE *out;
fpt *xsort, *ysort, bmax;
int index;
 int *order, count, j, i, bad;
int bestN, bestI;
 fpt den, bestDen;
#define CUTOFF ( bmax * ( xsort[order[i]] / xsort[order[j]] ) )
 if (!(order = (int *) UTL MEM_ALLOC( index *sizeof(int )))) return 0;
 for (i=0;i<index;i++) order[i]=i;</pre>
 bestN = bestI = bad = 0;
 bestDen = 0.0;
 fpt_heapsort(index, xsort, order);
 for (j=0;count=0, bad=0, j<index ;j++)
    if (xsort[order[j]] <= 0.0) continue;</pre>
    for (i=0; i<=j; i++)
       if (ysort[order[i]] <= CUTOFF) count++;</pre>
      else
                                     bad++;
                                                 */
     /* loop over all d <= this distance</pre>
    if ( (den = count/ bmax / xsort[order[j]] *2.0) > bestDen)
        {bestDen = den; bestI = j; bestN = index - bad;}
      /* loop over all distances
 den = bmax * xsort(order[index-1]);
 bestN,index,den-xsort[order[bestI]]*bmax/2.0, den);
 UBS OUTPUT MESSAGE(out, msg);
 UTL MEM FREE (order);
 return 1;
```

```
/* n is number of elements
   arrin is array of floats to be sorted
   indx is array of ints initially 0...n-1
*/
int fpt_heapsort(n,arrin,indx)
int n;
fpt *arrin;
int *indx;
int 1, ir, indxt, i, j;
fpt q;
1 = n/2 ;
ir = n - 1;
             /* the "10" loop */
while (TRUE)
  if (1>0) { indxt = indx[--1]; q = arrin[indxt]; }
  else
      indxt = indx[ir]; q = arrin[indxt];
      indx[ir--] = indx[0];
      if ( ir == 0 )
       { indx[0] = indxt; return 1; } /* <=== Only way out ! */
  i = 1;
  i = 1;
             +1;
  j = 1 + 1
  while (j <= ir) /* the "20" loop */
    indx[i] = indxt;
```

 $/\star$ SECTION III-A. Declarations for all non-standard data structures referenced in the C code functions shown in Sections I and II. $\star/$

```
**********
            Molecule and Supporting Structure Definitions
                 John McAlister
                                       09-Aug-1985
     This file contains the definitions for the molecular data struc-
     tures required within SYBYL. The contents of this file are des-
     described in detail in the document "SYBYL Molecular Data Struc-
           **********
/* Define the molecule descriptor template
                                                                        */
   typedef struct molecule struct {
                          /* pointer to molecule name
                                                                        */
     char
                *name;
                            /* molecule type
     i32
                type;
                            /* list of dictionaries used with molecule
     List Ptr
                 dict;
                            /* molecule status
     i32
                status;
                *comment;
                           /* pointer to comment for molecule
     char
                cre_time; /* creation time/user/version stamp
mod_time; /* modification time/user/version stamp
     stamp
     stamp
                 max_props; /* maximum properties currently allocated
     int
                            /* number of molecular properties
     int
                nprops;
                props; /* pointer to list of properties
max_feats; /* maximum features currently allocated
     List Ptr
     int
                nfeats; /* number of molecular features
     int
     List_Ptr
                            /* pointer to list of molecular features
                feats;
                 max_subst; /* maximum substructures currently allocated*/
     int
                         /* number of substructures in molecule
     int
                 nsubst;
     List Ptr
                            /* pointer to list of substructures
                subst;
                 subst_roots; /* pointer to list of root subst offsets
     List_Ptr
                 max_atoms; /* maximum atoms currently allocated
     int
                            /* number of atoms in molecule
     int
                natoms:
                atoms;
                           /* pointer to atom array segment list
     List Ptr
                 max_bonds; /* maximum bonds currently allocated
     int
                            /* number of bonds in molecule
     int
                 nbonds;
                            /* pointer to bond array segment list
/* type of atomic charges, if present
     List Ptr
                 bonds;
     int
                 charges;
                 vector[3]; /* translation vector for molecule
     fpt
                 matrix[9]; /* rotation matrix for molecule
     fpt
                 assoc_data; /* pointer to list of associated data
     List_Ptr
                                    descriptors
     } molecule, *mol_ptr;
      /* Define the atom entry record template
  typedef struct atom_struct {
                           /* atom name
     char
               *name;
     int
                           /* atom type
                type;
     i32
                status;
                            /* atom status
                            /* cumulative atom record number
     int
                recno;
                            /* atom id (logical atom number)
     int
                id;
                           /* link to next atom record
     int
                link;
                           /* offset to substructure containing atom
     int
               subst;
     List_Ptr
                            /\star pointer to list of properties for atom
               property;
     List Ptr
              feature;
                            /* pointer to list of features including
                            /*
                                this atom
                            /* number of bonds involving this atom
     int
                nbond;
```

31

	List_Ptr conn_atom; fpt xyz[3]; fpt charge; } atom, *atom_ptr;	<pre>/* pointer to list of bonded atoms /* coordinates of atom /* point charge on atom</pre>	*/ */ */
<pre>/* Define the atom array segment descriptor template typedef struct atom_seg_struct {</pre>			*/
	atom_ptr seg_head;	/* pointer to head of atom array segment	*/
	mol_ptr molecule;	/* pointer to molecule containing atom seg	*/
	int max_atom;	/* maximum number of atom records in seg	*/
	int natom;	/* number of filled atom records in seg	
	int used_atom;	/* offset to first filled record in segment	*/
		/* offset to first free record in segment	٠,
	<pre>} atom_seg, *aseg_ptr</pre>	;	
/*	Define the bond specifier	records pointed to by the atom records	*/
	typedef struct atom_conn	_struct { /* offset to target atom	*/
	int target;	/* offset to bond descriptor record	*/
			′
	} atom conn, *acon pt:	L;	

```
******* BOND DEFINITION ******
                                                                             */
/* Define the bond entry record template
   typedef struct bond_struct
                              /* bond type
                 type;
                              /* bond status
      i32
                 status;
                              /* cumulative bond record number
      int
                 recno;
                              /* bond id (logical bond number)
      int
                 id;
                              /* pointer to bond property list
/* pointer to list of features including
/* this bond
                 link;
      int
      List_Ptr
                 property;
      List_Ptr
                 feature;
                              /* offset to origin atom substructure
                 o subst;
      int
                              /* offset to atom at bond origin
      int
                 origin;
                              /* offset to target atom substructure
      int
                 t_subst;
                              /* offset to atom at bond destination
      int
                 target;
      } bond,
                *bond ptr;
/* Define the bond array segment descriptor template
                                                                             */
   typedef struct bond_seg_struct {
                              /* pointer to head of bond array segment
                                                                             */
                 seg head;
      bond ptr
                              /* pointer to molecule containing bond seg
                 molecule;
      mol_ptr
                                                                             */
                              /* maximum number of bonds in segment
                 max_bond;
      int
                              /* number of filled bond records in seg
      int
                 nbond;
                              /* offset to first filled record in segment
                 used bond;
      int
                             /* offset to first free record in segment
      int
                 free_bond;
      } bond_seg, *bseg_ptr;
```

```
/*****************
                                                                            */
/* =====
             comfa.h =====
/* Regions are the set of points at which energy evaluations are made
          in the CoMFA method of QSAR. A region is defined as the union */
.
/*
          of a set of 3D boxes (which may be a single point in the
           limit) and their associated attributes. Attributes needed for */
           CoMFA purposes are outlined below.
                                                                            */
   *************
#ifndef
                QSAR COMFA_DEFINITIONS
                QSAR COMFA DEFINITIONS 1
#define
                "ta_types.h"
#include
                               /* dummy atom id */
#define
                DUMMY 26
                    20
                               /* lone pair atom id */
                _{\rm LP}
#define
typedef enum {
  FDENGY UNKNOWN,
  FDENGY ELECT,
  FDENGY STERIC,
  FDENGY HOMO,
  FDENGY_LUMO,
 DOCK_ELECT,
DOCK_STA_NOHB,
  DOCK_STA_HBD,
  DOCK STA HBA,
  DOCK_STB_NOHB,
  DOCK_STB_HBD,
  DOCK STB HBA } FldEngyTyp;
typedef enum {
  FDHD_ORIGINAL,
  FDHD FFIT,
  FDHD XTERN,
  FDHD_FUNC,
  FDHD_USER,
  FDHD_USR_AVG,
FDHD_DOCK,
  FDHD_AVG,
  FDHD SIG,
  FDHD_MAX,
  FDHD_MIN,
  FDHD_COEFF,
  FDHD_AVG_X,
  FDHD SIG X,
  FDHD FLD X,
  FDHD RANGE,
  FDHD PLS XWT
  FDHD_PLS_XLOAD,
 FDHD_FAC_LOAD,
FDHD_FAC_COMM,
FDHD_FAC_ROTLOAD,
FDHD_SIMCA_LOAD,
  FDHD_SIMCA_MODEL,
  FDHD_SIMCA_DISCRIM,
  FDHD_HBD } FldHowTyp;
```

```
typedef struct {
                     /\star corner with lowest values for each axis
  fpt lo[3],
                    /* " " hi-est " " "
       hi[3],
       stepsize[3];
                    /* increment between points
                    /* derived as 1 + (hi-lo + epsilon) / stepsize
  int nstep[3],
                                                                       */
                     /* n = product of nstep[i]
      n;
                     /* SYBYL atom type, for steric energy computation
  int atom_type;
                                                                      */
                    /* elemental charge at point, for electrostatics
  fpt pt_charge;
                     /* weight[n] is applied in all computations,e.g=1
  fpt *weight;
  int avg_type;
                     /* box of 'scale', sphere, sphere x vdw, ...?
                                                                       */
  fpt avg_scale;
                    /* scale whose meaning derived from avg_type
                                                                       */
                    /*
  int arb,
                       arbitrary int for later use
                    /*
     *parb;
                                    pointer "
                } Box, *BoxPtr ;
typedef struct {
  char *filename;
                     /* name of the region's file (if any)
  int n_boxes;
                     /* number of boxes which make up the region
                     /* number of points in this region altogether
  int n_points ;
 BoxPtr box_array;
                     /* box_array[n_regions], each one a Box
  int n_refs ;
                     /* number of CURRENT references to this memory
  long when made;
                     /* creation stamp
               } Region, *RegionPtr;
typedef struct {
 char *reg_name;
                      /* name of the region's file (if any)
 char *fld_name;
                      /* name of this field's file (if any)
 RegionPtr reference; /* the region referenced by this field
 FldEngyTyp fld;
                       /* what type of field is referenced here
 int num_avgd;
                      /* number of fields averaged into this one
 int curr_iter;
                      /* number of iterations in current field fit run
 char *mol_id;
                      /* unspecified molecule id,
                          e.g. dbname/molname/alignname
                                                                         */
 int n points ;
                      /\star number of points in associated region
                      /* whether electrostatics are MISSING when>max st
 int zap_el;
                     /* largest permitted absolute value of energy
 fpt max value;
 fpt *field_value;
                      /* values at each point of the field
                     /* number of CURRENT references to this memory
 int n refs ;
                     /* creation stamp
 long when made;
 int vol_avg_type;
                        /* added these 4 items 1/30/89 DEP */
 fpt scale_vol_avg;
 int dielectric;
 int repulsive;
 FldHowTyp how made;
                           /* perry's way = 1 or old way = 0 */
    } Field, *FieldPtr ;
```

```
/* molecule dependent information solicited by QSAR table operations,
   passed into COMFA column field evaluations
typedef struct {
 boolean already_field; /* whether a field name exists (otherwise alignment) */
                         /* name of alignment;
 char
         *some name;
                                                  NIl align==use as is (!)
                                                                                */
                         /* name of steric
         *steric name;
                                                   field (if applicable)
                                                                                */
                         /* name of electrostatic field (if applicable)
         *elect name;
                         /* points to steric field in memory (when there)
 FieldPtr sfld_p;
 FieldPtr efld_p;
                         /* points to elect. field in memory (when there)
} ComfaMol, *ComfaMolPtr;
/* molecule-independent information for CoMFA evaluations */
typedef struct {
int vol_avg ;
fpt vol_scale ;
int fld_types ;
                       /* case for volume averaging: 0,1,2=none,box,sphere(0)*/
                       /* scale for volume averaging (1.0)
                       /* case for what fields: 0,1,2=both,steric,elect.(0)
 fpt steric max;
                       /* maximum steric energy (30)
                                                                               */
 int repulsive;
                       /* steric repulsive exponent - 12,10,or 8 (12)
                                                                               */
                       /* maximum electrostatic energy (30)
 fpt elect_max ;
                                                                               */
 int dielectric;
                       /* case for dielectric (AS FORCE FIELD TAILOR)
                       /* case to drop elect inside steric max: 0,1=T,F (1)
 int elect_out;
 char *region_name;
                       /* name of region used in the CoMFA computations
                                                                               */
 FieldPtr sweight_fld; /* points to MEMORY field for weighting steric PLS
 FieldPtr eweight_fld; /* points to MEMORY field for weighting elect. PLS
                            /* perry's way = 1 or old way = 0 */
 FldHowTyp how done;
                       /* include dummies and lone pairs in steric field
 int du_lp_steric;
                          calculations */
 int du_lp_elect;
                       /* include dummies and lone pairs in electrostatic
                          field calculations */
 int spare1;
                       /* As of 6.1comfa , this is TAILOR!COMFA!TRANSFORM*/
 int spare2;
                       /* INDICATOR SCALE among other things
} ComfaTop, *ComfaTopPtr;
```

#endif

Section III-B. Functional descriptions of external procedures. (Routines that simply return dynamic memory to the heap are not described.)

BOND_V_ERING - TRUE if bond is in an external ring.

BOND V IRING - TRUE if bond is in an internal (simple) ring.

QSAR_FIELD_EVAL_GETOFF - provides coordinates for field computation when "volume averaging" is being done.

QSAR_FIELD_VDWTAB - returns steric parameters for the computation of the field contribution from the probe atom and each of the molecule atoms.

SYB_AREA_GET_MOLECULE - returns the internal representation of the molecule in some area or "container", if such exists.

SYB_ATAB_ATOMIC_NUMBER - returns the atomic number of the specified atom type.

SYB_ATAB_ATOMIC_WEIGHT - returns the atomic weight of the specified atom type.

SYB_ATAB_HBOND_ACCEPT - returns TRUE if the specified atomic type is a hydrogen-bond accepting atom.

SYB_ATAB_VDW_RADII - returns the atomic radius of the specified atomic type.

SYB_ATOM_FIND_ID - returns the internal representation of an atom referenced by its atom ID number (Atom IDs are guaranteed to be continuous but the ID of any single atom may change as atoms are added or deleted.)

SYB_ATOM_FIND_REC - returns the internal representation of an atom referenced by its record ID number. (Atom record IDs are invariant but there may be "holes" in their sequence such that the largest record ID may be greater than the number of atoms.)

SYB_ATOM_FIND_SET - returns the bitset of atoms corresponding to a list of atoms.

SYB_BOND_FIND_REC - returns the internal representation of a bond referenced by its (invariant) record ID number.

SYB_BTAB_MNEM_TO_TYPE - converts an ASCII representation of a bond type to its internal representation.

SYB_EXPR_ANALYZE - parses a user-entered ASCII description of atoms (e.g., M2(<H>) for all hydrogen atoms within molecule M2) into internally valid representations of molecule and atoms.

SYB_HBOND_DONORS - returns the set of IDs for atoms which are hydrogen-bonding hydrogens.

TAILOR_STORE_IT_HERE - returns the current value of a user- (and SPL-) accessible variable.

TBL_ACCESS_INDEX_TO_COLNAME - converts a user-provided MSS column ID to a column name (name is guaranteed to be a unique identifier).

TBL_GRAB_COMPLETE_FPTS - done returning multiple (scalar) values in an MSS column to an array.

TBL_GRAB_GET_FPTS_INV - in a multiple value retrieval, returns the value corresponding to a user-provided row ID.

TBL_GRAB_INIT_FPTS - set up for returning multiple (scalar) values in an MSS column to an array.

UBS_OUTPUT_MESSAGE - equivalent to fprintf()

UIMS2_VAR_GET_TOKEN - returns the current value of a global SPL variable.

UIMS2_WRITE_ERROR - writes text to the error output stream.

UTL_FILE_FCLOSE, UTL_FILE_FOPEN - equivalent to fclose() and fopen().

UTL_LIST_RETRIEVE - returns the next element on a linked list.

UTL_MEM_ALLOC - equivalent to malloc().

UTL_SET_AND_INPLACE. makes the first set logically equivalent to the second set, with only those bits that are also 1 in the third set becoming 1 in the first set.

UTL_SET_CARDINALITY - returns the number of bits that are 1 in a particular bitset.

UTL_SET_CLEAR - sets all bits in the set to 0.

UTL_SET_COPY_INPLACE - makes the first set logically identical to the second.

UTL_SET_CREATE - creates and returns an empty set of requested size.

UTL_SET_DELETE - sets the specified bit to 0.

UTL_SET_DIFF_INPLACE - makes the first set logically equivalent to the second set, with all bits that are 1 in the third set becoming 0 in the first set.

UTL_SET_EMPTY - TRUE if all bits in the set are 0.

UTL_SET_INSERT - sets the requested bit to 1.

UTL_SET_MEMBER - returns TRUE if the requested set bit equals 1.

UTL_SET_NEXT - returns the identity of the next non-zero bit in a set

UTL_SET_OR_INPLACE - makes the first set logically equivalent to the second set, with all bits that are 1 in the third set becoming 1 in the first set.

UTL_STR_CMP_NOCASE - non-case sensitive version of strcmp().

APPENDIX "B"

```
/* CODE. This code implements a PHORE_LOC column type and calculates a single
cell value (the Hydrogen Bonding Fingerprint for a molecule) within the SYBYL
Molecular Spreadsheet. It is to be understood that other supporting code handles
user input, user output, and disk file I/O. */
/* data structure for PHORE_LOC column type */
typedef
   struct PHORE {
                        /* user name for DISCO feature file - default
       char *disco fn;
appears below */
                        /* internal flag if DISCO feature file loaded */
       int
            disco in;
                        /* user name for defining region file */
       char *region_fn;
                        /* internal reference to region when loaded */
       RegionPtr rgn;
                        /* number of extra lattice points (each direction)
       int nfuzz;
for each PHORE feature */
                        /* set length (must agree with rgn contents or EVAL
       int nbits;
fails) */
   } PHORE, *PPHORE;
/*+E:QSAR PROC EVAL PHORE LOC */
          /****
      int QSAR_PROC_EVAL_PHORE_LOC(tablename, row, colname)
/*
/*
                               (PHORE LOC == lattice bitset )
/*
     Dick Cramer 31-Jul-95
     This module generates bitsets whose cardinality is equal to
     lattice points x 2 (# of sitepoint classes. For each
.////////////
     instance of a pharmacophoric point in the molecule being
     processed, the geometrically nearest (1+m)^3 bits in the
     bitset will be set to 1 (where m is user supplied).
                                                                    */
     NOTE: this routine explicitly requires that sets begin after a
                                                                    */
*/
*/
          first element that is the set size!!!
      Inputs
                                                                    */
/*
      Outputs
/*
/*
      User Required Definition Files
/*-E*/
int QSAR PROC_EVAL_PHORE_LOC(tablename, row, colname)
        *tablename, *colname;
char
int
        row;
```

```
mol ptr
                  mol;
     PPHORE
                  phr;
     int
                  err, status, nvalid, mol area;
    char
                  *dum;
     set_ptr
               print, qsar_proc_calc_phore set();
    FILE *fp;
/* get the molecule */
    if ( !TBL_UTL_GET_MOLECULE(tablename, row, FALSE, &mol) )
       if ( UTL_ERROR IS SET() )
                                                                 {err=1; goto
error;}
      else return FALSE;
    }
/* get the user-provided input data */
    if ( !TBL_ATTR_FIND_COLUMN_A(tablename, colname, "PROC_SUPPORT", &dum,
                                     (int *)&phr) )
                                                                {err=3; goto
error;}
/* retrieve DISCO stuff if not yet present */
    if ( ! phr->disco_in) {
if (!phr->disco_fn) {err=1; goto error;}
/* set appropriate tallor value, then initialize DISCO */
     sprintf( str, "SETVAR TAILOR!DISCO!FILE %s", phr->disco fn );
     UIMS2_EXEC_COMMAND( str );
     UIMS2_EXEC_COMMAND( "DISCO INIT" );
     phr->disco_in = TRUE;
/* retrieve region if not yet present */
    if (!phr->rgn ) {
         if ( !phr->region_fn) {err=1; goto error;}
         if (!(phr->rgn = QSAR_REGION_RETRIEVE( phr->region_fn ) ))
{err=4;goto error;}
        if (phr->rgn->n boxes > 1 ) {
                 sprintf( str, "WARNING: Region %s has %d boxes. Only first
will be used.\n",
                          phr->region_fn, phr->rgn->n boxes );
                  UBS_OUTPUT_MESSAGE( stdout, str );
        phr->nbits = 2 * phr->rgn->n_points;
/* evaluate this result, first the DISCO call */
    if (!( print = qsar_proc_calc_phore_set( mol, phr, &nvalid )) ) {err=12;
goto error;}
/* go store both the bitset in the MSS "Cell_Support" and the number of bits
actually set in the "CELL", so there's something for the user to see */
if ( !TBL_ACCESS_X_PUT_VALUE(tablename, row, colname, "CELL_SUPPORT",
                                  (int *)&print) )
                                                             {err=11; goto error;}
```

```
if ( !TBL_ACCESS_X_PUT_VALUE(tablename, row, colname, "CELL",
                                (int *) &nvalid) )
                                                           {err=11; goto
error;}
    return TRUE;
error:
   sprintf (str, "QSAR_PROC_EVAL_PHORE_LOC (%d)", err);
UTL_ERROR_ADD_TRACE (str);
    return FALSE;
set ptr qsar_proc_calc_phore_set( mol, phr, nvalid )
/* creates actual bitset */
    mol ptr
                mol;
    PPHORE
                phr;
                *nvalid;
    int
  set_ptr anset = NIL, pset = NIL, SYB_FEAT_FIND_ID_SET();
  feat ptr featp, SYB_FEAT_FIND_REC();
              a, SYB_ATOM_FIND_REC();
       err, elem, sitebase, ci, xybase, boff, lt_base[3], lt_off[3], loff =
0, hioff = 0;
  fpt tmp;
  BoxPtr
                bxptr;
  line_ptr cdp;
    if (!( anset = UTL_SET_CREATE( phr->nbits ) )) {err = 1; goto error;}
    *nvalid = 0;
    if (phr->nfuzz) {
        loff -= phr->nfuzz / 2;
        hioff += (phr->nfuzz + 1) / 2;
    bxptr = phr->rqn->box array;
    xybase = bxptr->nstep[\overline{0}] * bxptr->nstep[1];
/* generate the DISCO sites for this molecule, which .. */
    UIMS2 EXEC COMMAND( "ECHO %DISCO_SITES()" );
/* .. become "FEATURES" + "dummy atoms" within SYBYL's molecule data
structure */
    pset = SYB FEAT FIND ID SET(mol, FEAT_V_LINE, 1, mol->nfeats);
    if (pset ) {
  elem = -1;
  while((elem = UTL SET NEXT(pset,elem)) != NO_MORE_ELEM) {
     if (!(featp = SYB_FEAT_FIND_REC (mol,elem))) goto error;
     if ((featp->name[1] == 'S') && (featp->name[2] == '_')) {
/* have an H-bonding feature, it must represent a line \overline{*}/
        sitebase = featp->name[0] == 'A' ? 0 : phr->rgn->n_points;
/* the dummy atom at the end of the line is our H-bonding Tocus */
```

```
cdp = (line ptr) featp->dataptr;
        if (!(a = SYB ATOM FIND REC (mol, cdp->positn)) ) {err=2; goto
error;}
       for (ci = 0; ci < 3; ci++) {
                tmp = (a->xyz[ci] - bxptr->lo[ci]) / bxptr->stepsize[ci];
                lt_base[ci] = (int) (tmp < 0.0 ? tmp - bxptr->stepsize[ci] :
tmp );
/* cycle through all points touched by this locus that are also within the
region */
        for (lt_off[0] = lt_base[0] + loff; lt_off[0] <= lt_base[0] + hioff;</pre>
lt_off[0]++)
        if (lt off[0] >= 0 && lt_off[0] < bxptr->nstep[0])
          for (lt_off(1) = lt_base(1) + loff; lt_off(1) <= lt_base(1) +
hioff; lt_off[1]++
          if (lt_off[1] >= 0 && lt_off[1] < bxptr->nstep[1])
             for (1t off[2] = 1t base[2] + loff; lt off[2] <= 1t base[2] +
hioff; lt off[2]++)
             if (lt_off[2] >= 0 && lt_off[2] < bxptr->nstep[2] ) {
                        boff = xybase * lt_off[2] +
                                 (bxptr \rightarrow nstep[0]) * lt_off[1] +
                                lt off(0) + sitebase;
                        UTL SET INSERT( anset, boff );
                        (*nvalid)++;
        }
     }
    UTL SET DESTROY( pset );
    } /* pset exists */
    return( anset );
error:
    sprintf (str, "qsar_proc_calc_phore_set(%d)", err);
    UTL_ERROR_ADD_TRACE (str);
    return FALSE;
}
   This file determines the recognition of site points in Sybyl/DISCO.
   See the SYBYL DISCO manual for detailed documentation. The defined types
are
     (1) HB: the QUERY is searched in the SEARCH mode, and all occurences
#
              are assigned DISCO features according to the remaining
#
              specifications -- the three ATOMS refer to the atom number
#
              in QUERY such that the feature is DIST from the first atom
              at bond ANGLE with the first and second atom at each of the
              TORSIONS formed by the site point and the three ATOMS in order.
              A sitepoint of NAME is added at these extension points,
              -- and -- the first atom is assigned a feature complimentary
```

```
to the extension point (such as HBD_CO_ and RHBD_CO_). (2) HBex:differs from HB in that the angles and torsions are replaced
              by two other arguments: whether lone pairs are part of the
              extension point placement, and which ATYPE (generally LP
              and/or H) determine the direction of the sitepoints.
              ATOMS SEARCH DIST ANGLE TORSIONS
#TYPE NAME
                                                     QUERY
#==== ====
              -----
               4 2 1 NoDup 2.9
                                   120 "0.0 180.0" HevC(Any)=O[f]
     DS 02C2
HB
                                 119 "0.0 180.0" O[f]HC(:Hev):Hev
     DS 03Car
               1 3 4 All 2.9
HB
                                 119 "0.0 180.0" O[f]C(:Hev):Hev
     DS_O3Car_
                1 2 3 All 2.9
     DS O3Car_
               1 3 4 NoDup 2.9
                                   119 "0.0 180.0" O[f]HC(=0)
HB
                                    119 "0.0 180.0" O[f]C(=0)
HB
     DS_03Car_ 1 2 3 NoDup 2.9
                                  120 "0.0 180.0" C(:O[f]):O[f]
     DS_O3Car_ 2 1 3 All 2.9
DS_O3C3_ 1 3 6 NoDup 2.9
HB
                                  117 "60 180 300"
O[f]HC(Any)(Any)C(Any)(Any)Any
     DS_N3C3_ 1 4 5 NoDup 2.9
DS_O2S_ 3 2 1 All 2.9 12
                                  110 "60 180 300" N[f]H2ZC{Z:C&!C=O&!C:Hev}
                            120 "0.0 180"
                                                AnyS (=0) (=0) NH
#TYPE NAME ATOMS
                      SEARCH DIST LP ATYPE Query
#==== ====
                      _____ ___
             =====
                                              ========
HBex DS 03C3 2 1 3 NoDup 2.9 YES "LP H"
O[f]HC(Any)(Any)Z{Z:Hev&!C(Any)(Any)Any}
HBex DS_O3C3_ 3 1 2 NoDup 2.9 YES "LP" HBex DS_N3C3_ 2 1 4 Nodup 2.9 "" "H"
                                             O[f](Z)Z{Z:C&!C=Het}
N[f]H2YaZ{Z:Hev&!C}{Ya:C&!C=0&!C:Hev}
HBex DS_N3C3_ 2 1 3 NoDup 2.9 YES "LP H" N[f]H(Ya)Ya{Ya:C&!C=O&!C:Hev}
             3 1 2 NoDup
                             2.9 YES "LP"
HBex DS N3C3
N[f](Ya)(Ya)Ya{Ya:C&!C=0&!C:Hev}
HBex DS_N2C2_
              2 1 3 NoDup
                              3.0
                                   YES "H LP" N[f]H=C
HBex DS N2C2
                                   YES "H LP" Any~N[f]=C
               1 2 3 NoDup
                              3.0
HBex DS N2C2
                              3.0 YES "LP"
               1 2 3 NoDup
                                              Any\sim N[r]=C[r]
               2 1 3 NoTriv 3.0 YES "LP H" N[1]H:C:C:N[f]:C:@1
HBex DS_N2N2_
HBex DS_N2N2_
                2 1 3 NoTriv 3.0 YES "LP H" N[1]H:C:C:N[f]:C:@1
                               3.0 YES "LP"
HBex DS_N2N2_
                3 2 1 NoDup
                                               C:N[f]:Hev
     DS_03S_
                                    128 "0.0 180.0"
hb
               3 2 1 NoDup 2.9
                                                          HevS=O[f]
     DS_O3S_
               4 2 1 All 2.9
                                     "0.0 180.0"
hb
                                 128
                                                       HevS(=O[f])=O[f]
                                 128 "0.0 180.0"
hb
     DS 03S
               4 2 1 All 2.9
                                                       HevS(~O[f])(~O[f])~O[f]
     DS_O3N_
                                 128 "0.0 180.0"
hb
               3 2 4 All 2.9
                                                       HevN(O[f])O[f]
     DS_O2N
                                    128 "0.0 180.0"
                                                          HevN(Hev)~O[f]
               4 2 1 NoDup 2.9
hbex DS_N2N2_
                3 2 1 NoDup
                               3.0 YES "LP"
                                                          N:N[f]:N
                3 1 2 All 2.9 128 "0.0 180.0"
     DS_03P_
hb
                                                       P(~0)(~0)(~0)(~0)
hb
     DS O3P
                3 1 2 All 2.9 128 "0.0 180.0"
                                                       P(~0)(~0)(~0)
    #CLASSNAMES# Acceptor_site Donor_Atom DL
AS_HO3C2_ 1 3 4 All 2.9 119 "0.0 180.0" O[f]HC(:Hev):Hev
     HB
                                   117 "60 180 300"
HB
O[f]HC(Any) (Any) C(Any) (Any) Any
     AS N3C3 1 4 7 NoDup 2.9
                                 110 "60 180 300"
N[f]H2C(Any)(Any)C(Any)(Any)Any
                                 110 "60 180 300"
     AS N3C3 1 5 8 NoDup 2.9
N[f]H3C(Any)(Any)C(Any)(Any)Any
#TYPE NAME
               ATOMS
                        SEARCH DIST LP ATYPE Ouerv
```

```
#==== ====
                 =====
                           HBex AS_HN2C2_ 2 1 3 NoDup
                                   3.0 "" "H" NHC(Any)=O[f]
HBex AS_HN2C2_ 3 2 1 NoDup
HBex AS_HN2C2_ 6 5 4 NoTriv
                   3 2 1 NoDup 3.0 YES "LP H" C:N[f]H:Hev 6 5 4 NoTriv 3.0 YES "LP" N[1]H:C:C:N[f]:C:@1
HBex AS HO3C3
                 2 1 3 NoDup
                                   2.9 YES "LP H"
O[f]HC(Any) (Any) Z{Z:Hev&!C(Any) (Any) Any}
                                   3.0 YES "LP H" HevN[f]H=C
HBex AS_HN2C2_ 3 2 4 Nodup
HBex AS_HN2C2_
                                   3.0 YES "LP" HevN[f]=C
                  1 2 3 Nodup
HBex AS_HN2C2_ 2 1 4 Nodup
HBex AS_N3C3_ 2 1 4 Nodup
                                  3.0 "" "H"
                                                  N[f]H2C(N)=N
                                  2.9 YES "LP H"
N[f]H2C(Any)(Any)Z{Z:Hev&!C(Any)(Any)Any}
HBex AS N3C3
                2 1 5 Nodup
                                  2.9 YES "LP H"
N[f]H3C(Any) (Any) Z{Z:Hev&!C(Any) (Any) Any}
HBex AS_N3C3_ 2 1 3 NoDup
                                 2.9
                                       YES "LP H" N[f]H(Ya)Ya{Ya:C&!C=O&!C:Hev}
                                 2.9 YES "LP H" N[f]H2(Ya)Ya{Ya:C&!C=O&!C:Hev}
HBex AS_N3C3_ 2 1 4 NoDup
HBEX AS_N3C3_ 2 1 4 NODUP
HBEX AS_N3C3_ 2 1 3 NODUP
HBEX AS_N3C3_ 3 1 2 NODUP
HBEX AS_HN2C2_ 2 1 3 NODUP
HBEX AS_HN2C2_ 2 1 4 NODUP
HBEX AS_HN2C2_ 2 1 3 NODUP
HBEX AS_HN2C2_ 1 3 NODUP
                                 2.9 YES "LP H" N[f]H(Ya)(Ya)Ya{Ya:C&!C=O&!C:Hev}
                                 2.9 YES "LP" N[f](Ya)(Ya)Ya{Ya:C&!C=O&!C:Hev}
                                   3.0 YES "H LP" N[f]H=C
                                        YES "LP" N[f]=C~Any
                                   3.0
                                        nn nHn
                                   3.0
                                                      N[f]H2Hev(:Hev):Hev
                                   3.0 ""
                                             "H"
                                                      N[f]HHev(:Hev):Hev
HBex AS_HN2C2_ 1 2 3 NoDup 3.0 "" "H"
HBex AS_HNS3_ 6 5 2 NoDup 3.0 "" "H"
HBex AS_HN4_ 2 1 3 NoDup -3.6 "" "C*"
                                   3.0 "" "H"
                                                      HNC=Any
                                                     AnyS(=0)(=0)N[f]H
                                                  N[f](Z)(Z)(Z)Z\{Z:C\&!C=O\&!C:Hev\}
N:N[f]:N
                                                               P(~0)(~0)(~0)(~0)
                                                               P(~0)(~0)(~0)
```

APPENDIX "C"

EXPERIMENTAL DATA SETS					
Data Set	No. Of Cpds	Structure, Activity			
1 Uehling	9	camptothecin, DNA fragmentation			
2 Strupczewski	34	benzisoxazoles, ip Behavioral			
3 Siddiqi	10	adenosines, Brain A1 binding			
4 Garratt1	10	tryptamines, melanophore binding			
5 Garratt2	14	tryptamines, melanophore binding			
6 Heyl	11	deltorphin, opioid receptor (DAMGO)			
7 Cristalli	32	adenosines, A2a agonists			
8 Stevenson	5	piperidines, NK1 antagonism			
9 Doherty	6	triarylbutenolides, endothelin-A antag.			
10 Penning	13	SC-41930 analogs, LTB4 antagonism			
11 Lewis	7	oxazolinediones, NK1 binding			
12 Krystek	30	sulfonamides, endothelin-A antagonism			
13 Yokoyamal	13	oxamic acids, T3 binding			
14 Yokoyama2	12	oxamic acids, T3 binding			
15 Svensson	13	benzindoles, 5-HTA agonism			
16 Tsutsumi	13	peptidyl heterocycles, endopeptidase inhib			
17 Chang	34	biphenyl sulfonamides, AT1 binding			
18 Rosowsky	10	trimetrexate analogs, DHFR inhibition			
19 Thompson	8	peptidomimetic, HIV-1 protease inhibition			
20 Depreux	26	naphthylethyl amides, melatonin displ.			

Literature References for Data Sets:

- Uehling, D.E., Nanthakamur, S.S., Croom, D., Emerson, D.L., Leitner, P.P.,
 Luzzio, M.J., et al., Synthesis, Topoisomerase I Inhibitory Activity, and in Vivo
 Evaluation of 11-Azacamptothecin Analogs. J. Med. Chem. 1995, 38, 1106 (Table 2, with R₂=Et; IC₅₀ data.
- Strupczewski, J.T., Bordeau, K.J., Chiang, Y., Glamkowski, E.J., Conway, P.G., et al. 3-[[(aryloxy)alkyl]piperidinyl]-1,2-Benzisoxazoles as D2/5-HT2 Antagonists with Potential Atypical Antipsychotic Activity: Antipsychotic Profile of Iloperidone

- (HP873). J. Med. Chem. 1995, 38, 1119. (Tables 2 and 3 with n=3, X=0; ED₅₀ for inhibition of apomorphine-induced climbing.)
- 3. Siddiqi, S.M., Jacobson, K.A., Esker, J.L., Olah, M.E., Ji, Xi.-duo., et al., Search for New Purine- and Ribose-Modified Adenosine Analogs as Selective Agonists and Antagonists at Adenosine Receptors. J. Med. Chem. 1995, 38, 1174. (Table 1, R₂=H; K₁(A1), values estimated from % displacement and stereoisomers averaged as needed.)
- 4. Garratt, P. J., Jones, R., Tocher, D. A., Sugden, D., Mapping the Melatonin Receptor. 3. Design and Synthesis of Melatonin Agonists and Antagonists Derived from 2-Phenyltryptamines. J. Med. Chem. 1995, 38, 1132. (Table 1 and Table 2).
- Garratt, P. J., Jones, R., Tocher, D. A., Sugden, D., Mapping the Melatonin Receptor. 3. Design and Synthesis of Melatonin Agonists and Antagonists Derived from 2-Phenyltryptamines. J. Med. Chem. 1995, 38, 1132. (Table 1 and Table 2).
- 6. Heyl, D.L., Dandabuthla, M., Kurtz, K.R., Mousigian, C. Opioid Receptor Binding Requirements for the &-Selective Peptide Deltorphin I: Phe³ Replacement with Ring-Substituted and Heterocyclic Amino Acids. J. Med. Chem. 1995, 38, 1242. (Table 1; binding K₁ to DAMGO.)
- 7. Cristalli, G., Camaioni, E., Vittori, S., Volpini, R., Borea, P.A., et al. 2-Aralkynyl and 2-Heteroalkynyl Derivatives of Adenosine-5'-N-ethyluronamide as Selective A2a Adenosine Receptor Agonists. J. Med. Chem. 1995, 38, 1462.
- 8. Stevenson, G.I., MacLeod, A.M., Huscroft, I., Cascieri, M.A., Sadowski, S., Baker, R. 4,4-Disubstituted Piperidines: A New Class of NK₁ Antagonist. J. Med.

- Chem. 1995, 38, 1264. (Table 1.)
- Doherty, A.M., Patt, W.C., Edmunds, J.J. Berryman, K.A., Reisdorph, B.R., et al.
 Discovery of a Novel Series of Orally Active Non-Peptide Endothelin-A (ET_A)
 Receptor-Selective Antagonists. J. Med. Chem. 1995, 38, 1259. (Table 3; IC₅₀ ET_A.)
- 10. Penning, T.D., Djuric, S.W., Miyashiro, J.M., Yu, S., Snyder, J.P., et al. Second-Generation Leukotriene B₄ Receptor Antagonists Related to SC-41930; Heterocyclic Replacement of the Methyl Ketone Pharmacophore. J. Med. Chem. 1995, 38, 858.

 (Table 1, all; LTB₄ receptor binding.)
- Lewis, R.T., MacLeod, A.M., Merchant, K.J. Kelleher, F., Sanderson, I., et al.
 Tryptophan-Derived NK1 Antagonists: Conformationally Constrained Heterocyclic
 Bioisosteres of the Ester Linkage. J. Med. Chem. 1995, 28, 923.
- 12. Krystek, S.R., Hunt, J.T., Stein, P.D., Stouch, T.R. 3D-QSAR of Sulfonamide Endothelin Inhibitors. J. Med. Chem. 1995, 38, 659.
- 13. Yokoyama, N., Walker, G.N., Main, A.J. Stanton, J.L. Morrissey, M., et al. Synthesis and SAR of Oxamic Acid and Acetic Acid Derivatives Related to L-Thyronine. J. Med. Chem. 1995, 38, 695.
- 14. Yokoyama, N., Walker, G.N., Main, A.J. Stanton, J.L. Morrissey, M., et al. Synthesis and SAR of Oxamic Acid and Acetic Acid Derivatives Related to L-Thyronine. J. Med. Chem. 1995, 38, 695.
- 15. Haadsma-Svensson, S.R., Svensson, K., Duncan, N., Smith, M.W., Lin, Ch.-H. C-9 and N-Substituted Analogs of cis-(3aR)-(-)-2,3,3a,4,5,9b-Hexahydro-3-propyl-1H-benz[e]indole-9-carboxamide: 5HT1A Receptor Agonists with Various Degrees of

- Metabolic Stability. J. Med. Chem. 1995, 38, 725.
- 16. Tsutsumi, S., Okonogi, T. Shibahara, S., Ohuchi, S., Hatsushiba, E., et al.,

 Synthesis and Structure Activity Relationships of Peptidyl @-Keto Heterocycles as

 Novel Inhibitors of Prolyl Endopeptidase. J. Med. Chem. 1994, 37, 3492. (Table 2,

 X=CH₂CH₂;IC₅₀.)
- 17. Chang, L.L., Ashton, W.T., Flanagan, K.L., Chen, Ts.-Bau., O'Malley, S.S., et al., Triazolinone Biphenylsulfonamides as Angiotensin II Receptor Antagonists with High Affinity for Both the AT₁ and AT₂ Subtypes. J. Med. Chem., 1994, 37, 4464. (Table 1, R³ = (2-C1)C₆H₅; AT₁ [rabbit aorta] IC₅₀.)
- 18. Rosowsky, A., Mota, C.E., Wright, J.E., Queener, S.F., 2,4-Diamino-5-chloroquinazoline Analogs of Trimetrexate and Piritrexim: Synthesis and Antifolate Activity. J. Med. Chem. 1994, 37, 4522. (Table 2; rat liver IC₅₀.)
- 19. Thompson, S.K., Murthy, K.H.M., Zhao, B., Winborne, E., Green, D.W., et al. Rational Design, Synthesis, and Crystallographic Analysis of a Hydroxyethylene-Based HIV-1 Protease Inhibitor Containing a Heterocyclic P1'-P2' Amide Bond Isostere. J. Med. Chem. 1994, 37, 3100. (Table 2, X-Boc; apparent K_i.)
- 20. Depreux, P., Lesieur, D., Mansour, H.A., Morgan, P., et al. Synthesis and Structure-Activity Relationships of Novel Naphthalenic and Bioisosteric Related Amidic Derivatives as Melatonin Receptor Ligands. J. Med. Chem. 1994, 37, 3231.

APPENDIX "D"

A list of 736 commercially available thiols broken down into 231 clusters based on topomeric CoMFA field descriptors along with the systematic name applicable to each. The 231 clusters are sorted by proposed name, first by the "root" structure, ie., the fragment attached immediately to the -SH, and then by the substitution pattern on that "root" substructure. The names describe topologically equivalent hydrocarbons, ie., structures in which all monovalent atoms are replaced by hydrogens and the other atoms by carbons.

01	Oluston	Chryst	Structural
	Cluster		
ID	Size	Root ·	=======================================
======	======		Simple
1	26	aryl	2,3,5-Me
144	1	aryl	
177	1	aryl	2,3,5-Me-4-Pr
163 ^C	1	aryl	2,3-(4-(2,3-Pr)5het)5het0
151	1	aryl	2,3-(4-Bu)5hetO-5-Me
33	5	aryl	2,3-Benzo
80	2	aryl	2,5-Me
192	1	aryl	2,5-Me-3-iPe
7	14	aryl	2,6-NoH-3(4/5)-Me
27	6	aryl	2,6-NoH-3-Ar
107	2	aryl	2-(2-Bz) PheEt-4, 5-Benzo
189	1	aryl	2-(3,5-Me)Ar-4,5-Benzo
141	1	aryl	2-(4-Et)PhePr
205	1	aryl	2-(4-Stilbenyl)Stilbenyl
188	1		2-5hetCH2-4,5-Benzo
56	3	aryl	
138	1	aryl	2-Ar-3,5-Me
190	1		2-Ar-4,5-(3,4-Et)Benzo
41	6 -	aryl	2-Ar-4,5-Benzo
152	1	aryl	2-Bz
16	9	aryl	2-Et
85	2	aryl	2-NoH-3-Et-5-Me
106	2	aryl	2-PheEt-4,5-Benzo
77	2	aryl	2-PhePr
142	1	aryl	2-R8
121	2	aryl	2-Stilbenyl
97	2	aryl	3,4-(3-Me)Benzo
218	1	aryl	3,4-(a,b) Inden0
164	1	aryl	3,4-(a,b,(8-Ar)IndenO)-6-Me
98	2		3,4-(a,b,(c-Me)IndenO)
99	3		3,4-(a,b-Naphtho)
157	1		3,4-Ar
58	3 2		3,4-Benzo-5-Me
100		aryl	3,4-Benzo-6-tBu
37	5	aryl	3,5-Me
180	1	aryl	3-(2,3-Benzo-4-Et)5het
199	1	aryl	3-(2,3-Benzo-5-Me)5het
182	1	aryl	3-(2-Me-3-5het-5-Et)5het
115	2	aryl	3-(3-5het) 5het
193	1	aryl	3-(3-Ar)5het-4-Me
67	3 2	aryl	3-Ar
129		aryl	3-Ar-4-(2-Me) 5hetCH2
46	4	aryl	3-Ar-5-Me
155	1	aryl	3-Bz
82	2	aryl	3-Bz-5,6-Benzo
10	16	aryl	3-Me

70	3	aryl	3-Naphth
73	3	aryl	3-Pr-4-sBu-6-Me
95	2	aryl	3-iPr
88	2	aryl	4-Ar
81	2	aryl	4-Bz
48	4	aryl	4-Et
2	23	aryl	4-Me
92	2	aryl	4-R9+
90	4	aryl	4-iBu
19	8	aryl	6-NoH
148 ^C	1	aryl	(adenosine)
228	ī	aryl	(fluorescein)
12	10	5het	Simple
50	4	5het	2,3-(a,b-Naphtho)
139	1	5het	2,3-5hetO-4-Me
89	2	5het	2,3-Ar
173	1	5het	2-(2,5-Et)Ar-3-Et
69	3	5het	2-(2-Me)Ar-3-(2-Me)PheEt
198	1	5het	2-(2-Me)Ar-3-R10
174	1	5het	2-(2-sBu)-3-Et
171	1	5het	2-(3,5-Me)Ar-3-5het
170	1	5het	2-(3,5-Me)Bz-3,4-Benzo
123	2	5het	2-(3-Et)Ar-3-Bz
22	7	5het	2-(4-Et)Ar
202	1	5het	2-(4-Et)Ar-4-(4-Me)Ar
122	2	5het	2-(4-iPr)Ar-3-Bz
197	1	5het	2-5hetCH2-3-(4-tBu)Ar
6	14	5het	2-Ar
225	1	5het	2-Ar-3-(2-Ar)5hetBu
224	1	5het	2-Ar-3-(2-Ar)5hetCH2
63	3	5het	2-Ar-3-(2-Bz)Ar
178	2	5het	2-Ar-3-(2-Me)5het
72	3	5het	2-Ar-3-(3,4-Et)Bz
40	5	5het	2-Ar-3-(3-Ar)5HetEt
183	1	5het	2-Ar-3-(3-Ar) PhePr
64	3	5het	2-Ar-3-(3-Ar-5-Me)5het
105	2	5het	2-Ar-3-(3-Me)Ar
160	1	5het	2-Ar-3-(4-Ar)Cyhx
146	1	5het	2-Ar-3-(4-Ar)CyhxCH2 2-Ar-3-(4-PheEt)Ar
203	1	5het	2-Ar-3-(4-FHEEC)Ar 2-Ar-3-(tBu)Ar
126	2	5het	2-Ar-3-(CBU/Ar 2-Ar-3-Ar
17_	9	5het	
211 ^C	1	5het	2-Ar-3-Benzylidene 2-Ar-3-IndenCH2
124	2	5het	
28b	6	5het	2-Ar-3-Me
30	6	5het	2-Ar-3-PhePr
204	1	5het	2-Ar-5-(4-(2,4-Me)Bz)Ar
79	2	5het	2-Bz
78	2 2	5het	2-Bz-3,4-Benzo
117		5het	
186	1 3	5het	
68		5het	2-Et 2-Et-3-(2-Me) PheEt
112	2	5het	Z-EC-J-(Z Me) FileDe

```
2-Me-3, 4-(3-Me)Benzo
           2
                     5het
128
                              2-Me-3, 4-Benzo
           2
                     5het
 93
                              2-Me-3-(2,3,4-Me) 5het
                     5het
           3
 61
                              2-Me-3-(2,3-Benzo-4-Et)5het
                     5het
           1
181
                              2-Me-3-(3-Ar)5het
                     5het
 49
           4
                              2-Me-3-(3-Ar) 5hetPr
           2
                      5het .
 86
                              2-Me-3-(3-Ar-5-Me) 5het
           2
                      5het
 91
                              2-Me-3-(3-Bz)Ar
          17
                      5het
  4
                              2-Me-3-(4-tBu)PheEt
                      5het
           1
172
                              2-Me-3-5Het
           5
                      5het
 38
                              2-Me-3-Me
          10
                      5het
 13
                              2-Me-3-Pe
                      5het
222
           1
                              2-Me-3-PheEt
           3
 66
                      5het
                              2-Me-3-PhePr
 29
           6
                      5het
                              2-Me-3-R8+
           3
                      5het
 71
                              .2-Me-5-Bu
           2
                      5het
108
                              2-Pe-3-Ar
           2
                      5het
127
                              2-Pr
           3
                      5het
 54
                              2-R12
            1
                      5het
221
                              2-iBu-3,4-iPe
                      5het
            1
187
                              2-iPe-3,4-Benzo
                      5het
143
            1
                              3,4-(2,4-Me)Benzo
            2
                      5het
 96
                              3,4-(3-Ar)Benzo
162
            1
                      5het
                              3,4-(3-Hx) Benzo
169
            1
                      5het
                              3,4-(3-Pr)Benzo
            2
                      5het
 94
                              3,4-(a,b-Napththo)
210
            1
                      5het
                              3,4-Benzo
           15
                      5het
 36
                              3-(2,4-Me)Bz
                      5het
176
            1
                      5het
                              3 - (3, 5 - Me) Ar
            1
196
                              3-(3-Ar)5het
                      5het
            1
159
                              3 - (3 - Bz) Ar
                      5het
 42
            4
                              3-(3-Me) PheEt
            1
                      5het
200
                              3 - (4 - Me) Ar
            2
                      5het
113
            2
                      5het
                              3-(4-tBu)Ar
125
                              3-(A1-4-Et) PheEt
                      5het
191
            1
                               3-(B-Ar)PhePr
145
            1
                      5het
                      5het
                               3-5hetCH2
114
            2
                               3-Ar
            8
                      5het
  18
                               3-Ar(2-thia)
            3
                       5het
  59
            3
                       5het
                               3-Bu
  65
                               3-Me-5-H
            7
                       5het
  24
                               3-Me-5-NoH
            6
                       5het
  44
                               3-Pe
            5
                       5het
 52
            2
                               3-PheEt
                       5het
 111
                               3-PhePr
            1
                       5het
 153
  32b
            6
                       5het
                               3-Pr
                               3-R13
                       5het
 223
            1
                               (chrysen0)
            1
                       5het
 185
            5
                     alkyl
                               Simple
  34
                               (3) (B1) (B1)
            2
 104
                      alkyl
            3
                      alkyl
                               (3-Me)PhePr
  62
                               (3:4)
                      alkyl
           18
   3
                               (3:4)(A1)
  14
            9
                      alkyl
```

```
(3:4)(B1)
                     alkyl
 60
           3
                               (4) (A1) (A-tBu) (C1) (C1)
            1
                     alkyl
226
                               (4) (D1) (D1)
            4
                     alkyl
 45
                               (4-Me) PhePr
            7
                     alkyl
 35
                               (4-iPe) PhePr
            1
                     alkyl
168
                     alkyl ·
                               (5) (A1)
            4
 47
                               (5) (B1) (E-(2-Ar-5-Me)5het)
            1
                     alkyl
179
            2
                     alkyl
                               (5) (B3)
103
                               (5)(C1)(C1)
            2
                     alkyl
 76
                               (5) (C2)
            2
                     alkyl
 83
                               (5) (C2) (D2) (D2)
            1
                     alkyl
216
                               (5:6)(D1/B1/F1)
                     alkyl
            8
 43
                               (5:7)
           15
                     alkyl
  5
                               (6) (B8) (C1) (E1) (E1)
                     alkyl
            1
158 -
                               (6) (F-Ar)
            1
                     alkyl
140
                              \cdot(7) (A8) (F1)
            1
                     alkyl
166
                               (7) (D3) (D3)
 53
            3
                     alkyl
                     alkyl
                               (8) (C3)
            1
207
                               (8:11)
                      alkyl
           13
  8
                               (9) (B4) (G3)
                      alkyl
            1
206
                               (10)(B1)(E5)(E1)
            3
                      alkyl
 75
                               (10)(C1)(E5)(E2)
            1
                      alkyl
136
                               (10+)(B1)
                      alkyl
            8
 20
                               (11+)(B1)
            7
                      alkyl
 39
154<sup>C</sup>
            1
                      alkyl
                                (12) (A-PheEt)
                                (12) (F6) (F1)
            1
                      alkyl
230
                                (12) (F6) (F6)
            2
                      alkyl
131
                                (12+)
            9
                      alkyl
 15
                                (13)(E4)
            1
                      alkyl
137
                                (A-Ar) (A-Ar) Bz
            1
                      alkyl
231
                                (A-Bz) (A-Bz) PheEt
229
            1
                      alkyl
                                (A1) PheEt
            1
                      alkyl
184
                                (cholesterol)
            1
227¢
                      alkyl
                                (cryptate)
 214<sup>C</sup>
            1
                      alkyl
             7
                                PheBu
                      alkyl
  23
                                PheEt
             3
                      alkyl
  74
                                PhePr
  25b
                      alkyl
             6
                                Simple.
  11
            10
                     benzyl
                                2,4,5-Me
             2
                     benzyl
 102
                                2,4,6-Me
             3
                     benzyl
  57
                                2-(3-(2-Et)Ar)Ar
             2
                     benzyl
 217
                                2-Et-3-(2,3-Et-5-Me)Ar-5-Me
             1
                     benzyl
 213
                                2-R8-3-Naphthyl-4,5-Benzo
             1
                     benzyl
 212
                                2/3-Me
            13
                     benzyl
   9
             2
                     benzyl
                                3,4-Benzo
  84
             2
                                3,5-Me
                     benzyl
 132
                                3-(4-Stilbenyl)Stilbenyl
             2
                     benzyl
 130
                                4-(3-Ar)Ar
             2
                     benzyl
 134
                                4-Et
             7
  21
                     benzyl
             6
                                4-Me
  26b
                     benzyl
             1
                     benzyl
                                4-PhePr
 156
             1
                                4-tBu
                     benzyl
 201
             2
                                Ar..(2-Et)Ar
                    alkenyl
 135
```

		- 111	λ - //-Pα\λ»
220	1	alkenyl	Ar(4-Bz)Ar
116	2	alkenyl	ArAr
133	2	alkenyl	ArBz
110	2	alkenyl	Et.CN.CONH2
87	2	alkenyl	NH2.CN.N=NPh
119	2	alkenyl	P(NMe2)3Ar
120	2	alkenyl	P(Pr)3Ar
118	2	alkenyl	P(iPe)3Ar
51	4	alkenyl	PCyhx3Ar
195 ^C	1	alkenyl	PEt3(2-Bz)Ar
31 ^b	6 •	alkenyl	PEt3Ar
194	1	alkenyl	PEt3Bz
109	2	alkenyl	PheEt.CN.CONH2
101	2 c	yclohexyl	Simple
149	1 c	yclohexyl	1-Me-2,4-CMe2
55	3 c	yclohexyl	2,3,4,5-iBu
147		yclohexyl	2,3,4-iBu-5-iPe
209	1 c	yclohexyl	2-(3,4-PheEt) 5het-6-Me
208		yclohexyl	2-Me-3,5-CMe2
167	1 c	yclohexyl	2-Me-4-sPe
165	1 c	yclohexyl	2-iPr-3,5-Me
150	1 c	yclohexyl	3-sPe-6-Me
161	1 c	yclohexyl	4-Et-4-iBu
219	1 c	yclohexyl	(complex)
175		clopentyl	2-Ar-4-spiro
215		clopentyl	3-PhePr

aTo generate these names, <u>all heteroatoms are first replaced by carbon</u> (to produce the simplest common topology) and a particular structure is chosen from among these topologies as the "most typical" of that cluster, if possible to contain the largest substructure that distinguishes that cluster from all others.

Within the name of a substitution, numbers indicate positions when substitution is on a ring, but chain length when substitution is on a chain (numbers separated by a colon indicate a range of chain lengths). Also, within a chain, letters indicate a position of substitution. (For example, (C2) describes a two atom branching from the third position of a chain, while 3-PhePr describes a phenyl propyl skeleton attached to the 3-position of a ring.)

A dot notation (.) separates the three possible substituents on an alkenyl root, the substituent order being same carbon as the -SH substituent, then the position *trans* to the -SH, and finally *cis* to -SH.

The above notwithstanding, <u>any</u> name enclosed completely in parentheses takes its usual structural meaning.

Here are structural descriptions for each name abbreviation in the above table, mostly in SLN (SYBYL Line Notation), listed alphabetically. (SLN extends SMILES with the following concepts, among others. Hydrogens are explicit. Ring openings and closures begin with a number enclosed by [] and end with the matching number preceded by @ Other SLN symbols used in these SLN definitions are: ~ = any bond; - = single bond (used here to provide a reference for [R]): = aromatic bond; ! = the SLN following (here in parentheses) is not allowed; [F] = no additional atoms may be attached to the preceding atom; [!R] = preceding bond may not be in a ring; [R] = preceding bond must be in a ring.)

5het = 5Het = C[1]:C:C:C:C:@1. alkenyl = C=C. alkyl = $C^{[R]C}$. aryl = Ar = Phe = Ph = C[1]:C:C:C:C:C@1. benzyl = $Bz = HSC-[!R]C^{R]C}$. Bu = C-[!R]C-[!R]C-[!R]C-[!R]C. cyclohexyl = $Cyhx = C[1](-l=)C^{C}C^{C}C^{C}$. cyclopentyl = $C[1]^{-(-l=)}C^{C}C^{C}C^{C}$. Et = C-[!R]C. inden = $C[1]:C(^{C}X^{[2]}):C(^{@2}):C:C@1$. iBu = C-[!R]C-[!R]C(-[!R]C)-[!R]C. iPe = C-[!R]C-[!R]C-[!R]C(-[!R]C)-[!R]C. Me = C. naphth = $C[1]:C(^{C}X^{[2]}):C(^{@2}):C:C:C@1$. NoH = $C^{(C)}C$. O denotes ring fusion, e.g., benzo fuses a 6-membered aromatic ring. Pe = C-[!R]C-[!R

r,